

PCT/IL 0 3 / 0 0 5 7 8  
10/519638  
07 AUG 2003

REC'D 25 AUG 2003

WIPO

PCT

PA 1036328

**THE UNITED STATES OF AMERICA**

**TO ALL TO WHOM THESE PRESENTS SHALL COME:**

**UNITED STATES DEPARTMENT OF COMMERCE**

**United States Patent and Trademark Office**

**July 08, 2003**

**THIS IS TO CERTIFY THAT ANNEXED HERETO IS A TRUE COPY FROM  
THE RECORDS OF THE UNITED STATES PATENT AND TRADEMARK  
OFFICE OF THOSE PAPERS OF THE BELOW IDENTIFIED PATENT  
APPLICATION THAT MET THE REQUIREMENTS TO BE GRANTED A  
FILING DATE UNDER 35 USC 111.**

**APPLICATION NUMBER: 10/193,136**

**FILING DATE: July 12, 2002**

**PRIORITY  
DOCUMENT**

SUBMITTED OR TRANSMITTED IN  
COMPLIANCE WITH RULE 17.1(a) OR (b)

**By Authority of the  
COMMISSIONER OF PATENTS AND TRADEMARKS**



*P. Swain*  
**P. SWAIN**

**Certifying Officer**

**BEST AVAILABLE COPY**

10/193,136 PCT/PTO 27-2004

# DIRECTOR OF THE PATENT AND TRADEMARK OFFICE

WASHINGTON D.C. 20231

Case Docket No. 02/23451

Sir:

Transmitted herewith for filing is the Patent Application of

Inventor : Eduardo N. MITRANI

FOR : METHOD AND DEVICE FOR INDUCING BIOLOGICAL PROCESSES BY MICRO-ORGANS

## Enclosed are:

- ☒ 21 sheets of drawings
- ☒ Assignment to: Yissum Research Development Company of the Hebrew University of Jerusalem
- ☒ Applicant is entitled to Small Entity Status under 37 CFR 1.9 and 37 CFR 1.27
- ☒ Floppy Disc with Sequence Listing
- ☒ Sequence Listing Statement: The sequence listing information recorded in computer readable form is identical to the written (on paper or compact disc) sequence listing

## THE FILING FEE HAS BEEN CALCULATED AS SHOWN BELOW:

FOR:	Column 1 No. FILED	Column 2 No. EXTRA	SMALL ENTITY		OTHER THAN A SMALL ENTITY	
			RATE	FEE	RATE	FEE
Basic Fee				\$ 370		\$ 740
Total Claims	167 - 20 =	147	147 x \$ 9	\$ 1323	x \$ 18	\$
Independent Claims	12 - 3 =	9	9 x \$ 42	\$ 378	x \$ 84	\$
<input checked="" type="checkbox"/> Recordal of Assignment \$40			+\$40	\$ 40	+\$40	\$
(If the difference in Column 1 is less than "0", enter "0" in Col. 2)			TOTAL	\$ 2111	TOTAL	\$

☒ Please charge my deposit account No. 50-1407 in the amount of \$2111.

A duplicate copy of this sheet is enclosed.

☒ The Commissioner is hereby authorized to charge payment of the following fees associated with this communication or credit any overpayment to Deposit Account No 50-1407. A duplicate copy of this sheet is enclosed.

☒ Any additional filing fees required under 37 CFR 1.16.

☒ Any patent application processing fees under 37 CFR 1.17.

☒ The Commissioner is hereby authorized to charge payment of the following fees during the pendency of this application or credit any overpayment to deposit Account No. 50-1407. A duplicate copy of this sheet is enclosed.

☒ Any patent application processing fees under 37 CFR 1.17

☒ Any filing fees under 37 CFR 1.16 for presentation of extra claims.

Respectfully,

*Sol Sheinbein*  
Sol Sheinbein

Registration No. 25,457

## APPLICATION FOR PATENT

5 Inventor: Eduardo N. Mitrani

10 Title: METHOD AND DEVICE FOR INDUCING BIOLOGICAL  
PROCESSES BY MICRO-ORGANS

15 This application is a Continuation-In-Part of U.S. Patent Application  
No. 10/009,520, filed June 22, 2000, which is a National Phase of  
PCT/IL00/00365, filed June 22, 2000, which claims the benefit of priority  
from U.S. Provisional Patent Application No. 60/140,748, filed June 25, 1999,  
the specifications of all of which are hereby incorporated by reference.

20

FIELD AND BACKGROUND OF THE INVENTION

The present invention relates to a method, extract and pharmaceutical  
composition for inducing angiogenesis in a tissue of a mammal, and to a  
25 device for the preparation and delivery of micro-organs (also referred to  
herein as micro-organ explants), into a mammal.

During the last few years numerous research studies have provided  
new insights into the molecular mechanisms which induce and regulate cell  
growth, and in particular, angiogenesis. The discovery of angiogenic growth  
30 factors such as vascular endothelial growth factor (VEGF), basic fibroblast  
growth factor (bFGF), angiopoietin and others, has led researchers to consider  
the use of these factors as agents for revascularization of ischemic tissue  
regions. Several different approaches utilizing either gene therapy or

recombinant protein technology have been attempted. Although preliminary results in animals were promising, clinical tests so far conducted, produced disappointing results (Ferrara and Alitalo, 1999 *Nature Medicine* 5(12): 1359-1364).

5       The lack of success at the clinical level can be attributed, at least in part, to the gene therapy or recombinant protein technology utilized in these experiments.

It has been shown that *in vivo* angiogenesis is effected and regulated by a complex and dynamic set of factors, including both stimulators and  
10 inhibitors (see Iruela-Arispe and Dvorak, 1997 *Thrombosis and Haemostasis* 78(1), 672-677, Gale and Yancopolous, 1999 *Genes and Development* 13, 1055-1066). In addition, it is thought that a long-term sustained stimulus is required to induce angiogenesis. Therefore, the current gene therapy and recombinant growth factors techniques, which do not address these issues,  
15 cannot produce the conditions necessary for promoting *in vivo* angiogenesis.

Recently, the inventor of the present invention have described a method for producing micro-organs which can be sustained outside the body in an autonomously functional state for extended periods of time. Such micro-organs, their preparation, preservation and some uses thereof are  
20 described, for example, in U.S. Patent No. 5,888,720; U.S. Patent Application No. 09/425,233, and in PCT/US98/00594, which are incorporated herein by reference.



SUMMARY OF THE INVENTION.

According to one aspect of the present invention, there is provided a method of inducing angiogenesis in a tissue of a first mammal, the method comprising the step of implanting at least one micro-organ within the tissue of  
5 the first mammal, said at least one micro-organ being for producing a plurality of angiogenic factors and thereby inducing angiogenesis.

According to an additional aspect of the present invention, said at least one micro-organ is derived from organ tissue of a second mammal.

According to an additional aspect of the present invention, the first  
10 mammal and said second mammal are a single individual mammal.

According to an additional aspect of the present invention, said organ is selected from the group consisting of a lung, a liver, a kidney, a muscle, a spleen a skin and a heart.

According to an additional aspect of the present invention, said at least  
15 one micro-organ includes two or more cell types.

According to an additional aspect of the present invention, the first mammal is a human being.

According to an additional aspect of the present invention, said at least one micro-organ is cultured outside the body for at least four hours prior to  
20 implantation within the tissue of the first mammal.

According to an additional aspect of the present invention, said at least one micro-organ is prepared so as to retain viability when implanted within the tissue of the first mammal.

According to an additional aspect of the present invention, said at least one micro-organ has dimensions, such that cells positioned deepest within said at least one micro-organ are at least about 80 - 100 microns and not more than about 225-375 microns away from a nearest surface of said at least one  
5 micro-organ.

According to an additional aspect of the present invention, each of said plurality of angiogenic factors posses a unique expression pattern within said at least one micro-organ.

According to an additional aspect of the present invention, at least a  
10 portion of cells of said at least one micro-organ include at least one exogenous polynucleotide sequence selected for regulating angiogenesis.

According to an additional aspect of the present invention, said at least one exogenous polynucleotide sequence is integrated into a genome of said at least a portion of said cells of said at least one micro-organ.

15 According to an additional aspect of the present invention, said at least one exogenous polynucleotide sequence is designed for regulating expression of at least one angiogenic factor of said plurality of angiogenic factors.

According to an additional aspect of the present invention, said at least one exogenous polynucleotide sequence includes an enhancer or a suppresser  
20 sequence.

According to an additional aspect of the present invention, an expression product of said at least one exogenous polynucleotide sequence is

capable of regulating the expression of at least one angiogenic factor of said plurality of angiogenic factors.

According to an additional aspect of the present invention, said at least one exogenous polynucleotide sequence encodes at least one recombinant  
5 angiogenic factor.

According to another aspect of the present invention, there is provided a method of inducing angiogenesis in a tissue of a first mammal, the method comprising steps of:

- (a) extracting soluble molecules from  
10 at least one micro-organ; and
- (b) administering at least one predetermined dose of said soluble molecules extracted in step (a) into the tissue of the first mammal.

According to an additional aspect of the present invention, said soluble molecules are mixed with a pharmaceutically acceptable carrier prior to step  
15 (b).

According to an additional aspect of the present invention, said at least one micro-organ is derived from organ tissue of a second mammal.

According to an additional aspect of the present invention, said at least one micro-organ is cultured at least four hours prior to extraction of said  
20 soluble molecules.

According to an additional aspect of the present invention, said at least one micro-organ has dimensions, such that cells positioned deepest within said at least one micro-organ are at least about 80 - 100 microns and not more

than about 225-375 microns away from a nearest surface of said at least one micro-organ.

According to another aspect of the present invention, there is provided a pharmaceutical composition comprising, as an active ingredient, a soluble  
5 molecule extract from at least one micro-organ and a pharmaceutically acceptable carrier.

According to another aspect of the present invention, there is provided a micro-organ comprising a plurality of cells, wherein at least a portion of said plurality of said cells including at least one exogenous polynucleotide  
10 sequence, said at least one exogenous polynucleotide sequence being capable of regulating expression of at least one angiogenic factor expressed in said cells.

According to an additional aspect of the present invention, the micro-organ is derived from organ tissue of a second mammal.

15 According to an additional aspect of the present invention, the first mammal and said second mammal are a single individual mammal.

According to an additional aspect of the present invention, said organ is selected from the group consisting of a lung, a liver, other gut derived organs, a kidney, a spleen and a heart.

20 According to an additional aspect of the present invention, said at least one micro-organ includes two or more cell types.

According to an additional aspect of the present invention, the micro-organ has dimensions, such that cells positioned deepest within the micro-

organ are at least about 80 - 100 microns and not more than about 225 - 375 microns away from a nearest surface of the micro-organ.

According to an additional aspect of the present invention, said at least one exogenous polynucleotide sequence is integrated into a genome of said at  
5 least a portion of said plurality of said cells.

According to an additional aspect of the present invention, said at least one exogenous polynucleotide sequence includes an enhancer or a suppressor sequence.

According to an additional aspect of the present invention, an  
10 expression product of said at least one exogenous polynucleotide sequence is capable of regulating the expression of said at least one angiogenic factor.

According to another aspect of the present invention, there is provided a method of inducing angiogenesis in a tissue of a first mammal, the method comprising the steps of:

15 culturing at least one micro-organ in a growth medium to thereby generate a conditioned medium;

collecting said conditioned medium following at least one predetermined time period  
20 of culturing; and

administering at least one predetermined dose of said conditioned medium collected in

step (b) into the tissue of the first mammal to  
thereby induce angiogenesis in the tissue.

According to an additional aspect of the present  
invention, said at least one micro-organ is

5 derived from organ tissue of a second mammal.

According to an additional aspect of the present invention, said at least  
one micro-organ is cultured at least four hours prior to collection of said  
conditioned medium.

According to an additional aspect of the present invention, said at least  
10 one micro-organ has dimensions, such that cells positioned deepest within  
said at least one micro-organ are at least about 80 - 100 microns and not more  
than about 225-375 microns away from a nearest surface of said at least one  
micro-organ.

According to an additional aspect of the present invention, said  
15 growth medium is a minimal essential medium.

According to another aspect of the present invention there is  
provided apparatus for generating micro-organs from a tissue biopsy and  
for implanting the micro-organs into a subject, the apparatus comprising:

(a) a cutting chamber for cutting the tissue biopsy into a plurality  
20 of micro-organs; and

(b) an implanting mechanism for implanting the plurality of  
micro-organs into the subject, said implanting mechanism being operably  
coupled to said cutting chamber.

According to an additional aspect of the present invention, said cutting chamber has an inlet/outlet for introducing and removing reagents.

According to an additional aspect of the present invention, said cutting chamber has an inlet for introducing the tissue biopsy therein.

5 According to an additional aspect of the present invention, said apparatus comprises a viability testing chamber operably coupled to said cutting chamber for testing a viability of at least one sacrificial micro-organ of said plurality of micro-organs.

10 According to an additional aspect of the present invention, said implanting mechanism comprises a multi-channel planter and corresponding advancing elements for advancing said plurality of micro-organs from said cutting chamber to said multi-channel planter and further for implanting the plurality of micro-organs into the subject.

15 According to an additional aspect of the present invention, said apparatus comprises a processing chamber being operably coupled to said cutting chamber and said implanting mechanism for processing said micro-organs prior to said implanting.

20 According to an additional aspect of the present invention, said processing chamber has an inlet/outlet for introducing and removing processing reagents.

According to an additional aspect of the present invention, said cutting chamber comprises a cutting mechanism having a plurality of blades movable to cut the tissue biopsy into said plurality of micro-organs.

According to an additional aspect of the present invention, said blades are so disposed with respect to one another such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80-100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ.

According to an additional aspect of the present invention, said plurality of blades has a translatable angled cutting edge.

According to an alternative aspect of the present invention, each of said plurality of blades is a rotatable disc-blade.

According to another aspect of the present invention there is provided apparatus for generating micro-organs from a tissue biopsy, the apparatus comprising:

(a) a cutting chamber for cutting the tissue biopsy into a plurality of micro-organs; and

(b) a viability testing chamber operably coupled to said cutting chamber for testing a viability of at least one sacrificial micro-organ of said plurality of micro-organs.

According to an additional aspect of the present invention, said cutting chamber has an inlet/outlet for introducing and removing reagents.

According to an additional aspect of the present invention, said cutting chamber has an inlet for introducing the tissue biopsy therein.



According to an additional aspect of the present invention, said cutting chamber comprises a cutting mechanism having a plurality of blades movable to cut the tissue biopsy into said plurality of micro-organs.

According to an additional aspect of the present invention, said  
5 blades are so disposed with respect to one another such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225 - 375 microns away from a nearest surface of said micro-organ.

10 According to an additional aspect of the present invention, each of said plurality of blades has a translatable angled cutting edge.

According to an alternative aspect of the present invention, each of said plurality of blades is a rotatable disc-blade.

According to another aspect of the present invention there is  
15 provided apparatus for generating micro-organs from a tissue biopsy, the apparatus comprising:

(a) a cutting chamber for cutting the tissue biopsy into a plurality of micro-organs;

(b) a processing chamber being operably coupled to said cutting  
20 chamber for processing said micro-organs;

(c) an advancing mechanism for advancing said micro-organs from said cutting chamber into said processing chamber.

According to an additional aspect of the present invention, said processing chamber has an inlet/outlet for introducing and removing processing reagents.

According to an additional aspect of the present invention, said  
5 cutting chamber has an inlet/outlet for introducing and removing reagents.

According to an additional aspect of the present invention, said cutting chamber has an inlet for introducing the tissue biopsy therein.

According to an additional aspect of the present invention, said cutting chamber comprises a cutting mechanism having a plurality of  
10 blades movable to cut the tissue biopsy into said plurality of micro-organs.

According to an additional aspect of the present invention, said blades are so disposed with respect to one another such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of  
15 micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ.

According to an additional aspect of the present invention, each of said plurality of blades has a translatable angled cutting edge.

According to an alternative aspect of the present invention, each of  
20 said plurality of blades is a rotatable disc-blade.

According to another aspect of the present invention there is provided a method of generating micro-organs from a tissue biopsy and for implanting the micro-organs into a subject, the method comprising:

providing an apparatus which comprises:

(a) a cutting chamber for cutting the tissue biopsy into a plurality of micro-organs; and

(b) an implanting mechanism for implanting the plurality of  
5 micro-organs into the subject, said implanting mechanism being operably coupled to said cutting chamber.

placing the tissue biopsy in said cutting chamber and cutting the tissue biopsy into the plurality of micro-organs; and

using said implanting mechanism for implanting the plurality of  
10 micro-organs into the subject.

According to an additional aspect of the present invention, the micro-organs serve as angiopumps.

According to an additional aspect of the present invention, said cutting chamber has an inlet/outlet for introducing and removing reagents,  
15 the method further comprising washing said micro-organs in said cutting chamber prior to using said implanting mechanism for implanting the plurality of micro-organs into the subject.

According to an additional aspect of the present invention, said cutting chamber has an inlet for introducing the tissue biopsy therein, the  
20 method comprising placing the tissue biopsy in said cutting chamber through said inlet.

According to an additional aspect of the present invention, said apparatus further comprises a viability testing chamber operably coupled to

said cutting chamber for testing a viability of at least one sacrificial micro-organ of said plurality of micro-organs, the method further comprising testing said viability of said at least one sacrificial micro-organ of said plurality of micro-organs prior to using said implanting mechanism for  
5 implanting the plurality of micro-organs into the subject.

According to an additional aspect of the present invention, said implanting mechanism comprises a multi-channel implanter and corresponding advancing elements for advancing said plurality of micro-organs from said cutting chamber to said multi-channel implanter and  
10 further for implanting the plurality of micro-organs into the subject, the method comprising implanting the plurality of micro-organs into the subject using said advancing elements.

According to an additional aspect of the present invention, said apparatus comprises a processing chamber being operably coupled to said  
15 cutting chamber and said implanting mechanism for processing said micro-organs prior to said implanting, the method further comprising processing said micro-organs prior to said implanting.

According to an additional aspect of the present invention, said processing said micro-organs prior to said implanting comprises at least one  
20 a process selected from the group consisting of washing, transforming, culturing, and a combination thereof.

According to an additional aspect of the present invention, said processing said micro-organs prior to said implanting comprises culturing for at least one hour.

According to an additional aspect of the present invention, said  
5 processing said micro-organs prior to said implanting comprises transforming by introducing to at least a portion of cells of said micro-organs at least one exogenous polynucleotide sequence selected for regulating angiogenesis.

According to an additional aspect of the present invention, said at  
10 least one exogenous polynucleotide sequence is integrated into a genome of said at least said portion of said cells of said micro-organs.

According to an additional aspect of the present invention, said at least one exogenous polynucleotide sequence is designed for regulating expression of at least one angiogenic factor of said plurality of angiogenic  
15 factors.

According to an additional aspect of the present invention, said at least one exogenous polynucleotide sequence includes an enhancer or a suppresser sequence.

According to an additional aspect of the present invention, said at  
20 least one exogenous polynucleotide sequence is capable of regulating the expression of at least one angiogenic factor of said plurality of angiogenic factors.

According to an additional aspect of the present invention, said at least one exogenous polynucleotide sequence encodes at least one recombinant angiogenic factor.

According to an additional aspect of the present invention, said  
5 processing chamber has an inlet/outlet for introducing and removing processing reagents, the method comprising introducing at least one processing reagent into said processing chamber through said inlet/outlet.

According to an additional aspect of the present invention, said cutting chamber comprises a cutting mechanism having a plurality of  
10 blades movable to cut the tissue biopsy into said plurality of micro-organs, the method comprising using said plurality of blades to cut the tissue biopsy into said plurality of micro-organs.

According to an additional aspect of the present invention, said cutting chamber is designed and constructed such that once the tissue  
15 biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ, the method further comprising using said cutting chamber to cut the tissue biopsy into  
20 said plurality of micro-organs each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from said nearest surface of said micro-organ.

According to an additional aspect of the present invention, each of said plurality of blades has a translatable angled cutting edge, the method comprising translating said angled cutting edge with respect to the tissue biopsy, so as to cut the tissue biopsy into said plurality of micro-organs.

5 According to an alternative aspect of the present invention, each of said plurality of blades is a rotatable disc-blade, the method comprising moving said rotatable disc-blade with respect to the tissue biopsy, so as to cut the tissue biopsy into said plurality of micro-organs.

10 According to an additional aspect of the present invention, said tissue biopsy is derived from a tissue or organ selected from the group consisting of lung, liver, kidney, muscle, spleen, skin, heart, lymph node and bone marrow.

According to an additional aspect of the present invention, a donor of the tissue biopsy and the subject are the same individual.

15 According to an alternative aspect of the present invention, a donor of the tissue biopsy and the subject are different individuals.

According to an additional aspect of the present invention, a donor of the tissue biopsy is a human.

20 According to an alternative aspect of the present invention, a donor of the tissue biopsy is a non-human mammal.

According to an additional aspect of the present invention, said subject is a non-human mammal.

According to an alternative aspect of the present invention, said subject is a human.

According to an additional aspect of the present invention, said implanting the plurality of micro-organs into the subject is effected via  
5 transmucosal or parenteral administration routes.

According to an additional aspect of the present invention, said transmucosal or parenteral administration routes are selected from the group consisting of intramuscular, subcutaneous, intramedullary, intrathecal, direct intraventricular, intravenous, intraperitoneal, intranasal  
10 and intraocular administration routes.

According to a preferred aspect of the present invention, there is provided a device for micro-organ preparation and delivery, comprising:

a tissue scraper, for obtaining a tissue biopsy;  
a tissue cutter, for cutting the tissue biopsy into a plurality of  
15 fragments, forming a plurality of micro-organs; and

at least one implanting device, detachably coupled to said tissue cutter, for receiving a micro-organ, of said plurality of micro-organs, when coupled to said tissue cutter, and for implanting said micro-organ into a subject, after decoupling from said tissue cutter.

20 According to an additional aspect of the present invention, said device is sealed within a base, a ramp, and a casing.

According to an additional aspect of the present invention, said device includes a control system.



According to an additional aspect of the present invention, said device includes at least one automated travel mechanism for transferring the tissue biopsy from one region of said device to another.

According to an additional aspect of the present invention, said  
5 tissue scraper is adapted for scraping said tissue to a predetermined width.

According to an additional aspect of the present invention, said tissue scraper is adapted for scraping said tissue to a predetermined length.

According to an additional aspect of the present invention, said tissue scraper is adapted for scraping said tissue to a predetermined  
10 thickness.

According to an additional aspect of the present invention, said tissue scraper has a replaceable blade.

According to an additional aspect of the present invention, said device includes a washing apparatus for rinsing the tissue biopsy.

According to an additional aspect of the present invention, said  
15 washing apparatus is operative for applying a medium to the tissue biopsy.

According to an additional aspect of the present invention, said device is further operative as a tissue treatment chamber.

According to an additional aspect of the present invention, said  
20 device includes apparatus for controlling the temperature therein.

According to an additional aspect of the present invention, said tissue cutter comprises a plurality of parallel, surgical-grade blades, designed to cut the tissue biopsy into said plurality of fragments, forming said micro-

organs, such that cells positioned deepest within any one of said micro-organs are at least about 80 - 100 microns and not more than about 225 - 375 microns away from a nearest surface.

According to an additional aspect of the present invention, said tissue  
5 cutter comprises a plurality of parallel surgical-grade blades, arranged at an angle to the tissue biopsy.

According to an alternative aspect of the present invention, said tissue cutter comprises a plurality of parallel surgical-grade blades, arranged as rotatable disc-blades.

10 According to an additional aspect of the present invention, said device comprises a viability testing chamber for testing a viability of at least one micro-organ of said plurality of micro-organs.

According to an additional aspect of the present invention, said tissue cutter is operative to cut the tissue biopsy, to form said micro-organs,  
15 and to arrange each of said micro-organs on a single micro-organ guide of a plurality of micro-organ guides, in a single operation.

According to an additional aspect of the present invention, said at least one implanting device includes a slim housing, adapted for percutaneous insertion, and operable to receive one of said plurality of  
20 micro-organ guides.

According to an additional aspect of the present invention, said at least one implanting device includes a plurality of implanting devices, each operable to receive one of said plurality of micro-organ guides.

According to an additional aspect of the present invention, each of said plurality of micro-organ guides includes a position marker for indicating when said micro-organ, arranged on it, is positioned for implanting.

5        According to an additional aspect of the present invention, each of said micro-organ guides includes a notch for breaking off a distal portion thereof, to allow said micro-organ, arranged on it, to form a leading edge.

      According to an additional aspect of the present invention, each of said plurality of micro-organ guides includes a position marker for  
10        indicating when said micro-organ, arranged on it, is implanted.

      According to an additional aspect of the present invention, said device is disposable.

      According to another preferred aspect of the present invention, there is provided a method for micro-organ preparation and delivery, comprising:

15        scraping a tissue biopsy;

      cutting the tissue biopsy to a plurality of fragments, forming a plurality of micro-organs; and

      implanting at least one of said plurality of one micro-organs.

      According to another preferred aspect of the present invention, there  
20        is provided a method for micro-organ preparation and delivery, comprising:

      employing a device for micro-organ preparation and delivery, which includes:

      a tissue scraper, for obtaining a tissue biopsy;

a tissue cutter, for cutting the tissue biopsy into a plurality of fragments, forming a plurality of micro-organs; and

at least one implanting device, detachably coupled to said tissue cutter, for receiving a micro-organ, of said plurality of micro-organs, when coupled to said tissue cutter, and for implanting said micro-organ into a subject, after decoupling from said tissue cutter;

scraping the tissue biopsy, with said tissue scraper;

cutting the tissue biopsy to said plurality of fragments, forming said plurality of micro-organs, with said tissue cutter;

10 mounting said micro-organ, of said plurality of micro-organs, on said at least one implanting device;

decoupling said at least one implanting device; and

implanting said micro-organ, with said at least one implanting device.

15 According to an additional aspect of the present invention, the micro-organ serves as an angiopump.

According to an additional aspect of the present invention, said method includes treating the tissue biopsy, prior to implanting.

According to an additional aspect of the present invention, said  
20 treating is selected from the group consisting of washing, transforming, culturing, and a combination thereof.

According to an additional aspect of the present invention, said cutting includes arranging each of said micro-organs on a single micro-organ guide of a plurality of micro-organ guides.

According to an additional aspect of the present invention, said  
5 method includes disposing said device after a single use

According to an additional aspect of the present invention,  
said cutting further includes cutting to a first plurality of tissue  
fragments, forming a first plurality of micro-organs; and

said implanting further includes implanting a second plurality of  
10 micro-organs, wherein said second plurality is selected from the group  
consisting of a plurality which is equal to said first plurality, a plurality  
which is smaller than said second plurality by one, and a plurality which is  
smaller than said second plurality by two.

According to an alternative aspect of the present invention,  
15 said cutting further includes cutting to a first plurality of tissue  
fragments, forming a first plurality of micro-organs; and

said implanting further includes implanting a second plurality of  
fragments, wherein said second plurality is smaller than said first plurality  
by one,

20 wherein said method further includes using an edge fragment for a  
viability test.

According to an alternative aspect of the present invention,

said cutting further includes cutting to a first plurality of tissue fragments, forming a first plurality of micro-organs; and

said implanting further includes implanting a second plurality of tissue fragments, wherein said second plurality is smaller than said first plurality by two;

wherein said method further includes using a first edge fragment for a viability test; and

discarding a second edge fragment.

According to an additional aspect of the present invention, said cutting includes cutting the tissue biopsy into said plurality of fragments, forming said micro-organs, such that cells positioned deepest within any one of said micro-organs are at least about 80 microns and not more than about 375 microns away from a nearest surface.

According to an additional aspect of the present invention, said cutting includes cutting the tissue biopsy into said plurality of fragments, forming said micro-organs, such that cells positioned deepest within any one of said micro-organs are at least about 100 microns and not more than about 225 microns away from a nearest surface.

According to an additional aspect of the present invention, said implanting further includes implanting a plurality of micro-organs within a preselected area of said subject, for a predetermined area concentration of micro-organs.

According to an additional aspect of the present invention, said implanting further includes implanting a plurality of micro-organs within a preselected volume of said subject, for a predetermined volume concentration of micro-organs.

5 According to an additional aspect of the present invention, said tissue biopsy is a split-thickness tissue biopsy.

The present invention successfully addresses the shortcomings of the presently known configurations by providing a method, extract, and pharmaceutical composition for inducing angiogenesis in a tissue of a  
10 mammal, and a device for the preparation and delivery of micro-organs into a mammal.

#### BRIEF DESCRIPTION OF THE DRAWINGS

The invention is herein described, by way of example only, with  
15 reference to the accompanying drawings. With specific reference now to the drawings in detail, it is stressed that the particulars shown are by way of example and for purposes of illustrative discussion of the preferred embodiments of the present invention only, and are presented in the cause of providing what is believed to be the most useful and readily understood  
20 description of the principles and conceptual aspects of the invention. In this regard, no attempt is made to show structural details of the invention in more detail than is necessary for a fundamental understanding of the invention, the

description taken with the drawings making apparent to those skilled in the art how the several forms of the invention may be embodied in practice.

In the drawings:

FIG. 1 is a photograph showing neo-vascularization around an  
5 implanted micro-organ (marked with arrow);

FIG. 2 is a graph illustrating the relative levels of various angiogenic factors expressed in transplanted micro-organs. Ang1 - angiopoietin 1, Ang2 - angiopoietin 2, MEF2C - myocyte enhancer factor 2C, VEGF - vascular endothelial growth factor;

10 FIG. 3 is an angiogenic factor-specific RT-PCR of RNA extracted from micro-organs cultured outside the body for various time points following preparation. Actin - beta-actin (control);

FIG. 4 is a graph representing semi-quantitative data obtained by densitometry of the RT-PCR products shown in Figure 3, normalized to the  
15 intensity of the beta-actin RT-PCR product (control);

FIG. 5 is a histogram representing the gating pattern of common iliac-ligated rats implanted with micro-organs or sham implanted (control). (n) = 13. P values for the three time groups (from left to right) are 0.16, 1 and 0.841. Scores: 0-full functionality 9-total inability to move the limb, 10 loss  
20 of the limb;

FIG. 6 is a histogram representing the same experimental group as in Figure 5 with the exception that the animals were now exerted prior to



scoring gating behavior. P values for the three time groups are (from left to right) 0.0001, 0.0069 and 0.06;

FIG. 7 is a histogram representing the gating pattern of common iliac-ligated mice implanted with micro-organs or sham implanted. Scores: 0-full  
5 functionality 9-total inability to move the limb, 10 loss of the limb. P values for the three time groups are (from left to right) 0.00025, 0.00571 and 0.07362;

FIG. 8 is an image illustrating a mouse spleen derived micro-organ (marked with MC arrow) six months following implantation into a  
10 subcutaneous region of a syngeneic mouse. One of the newly formed blood vessels surrounding the micro-organ is marked with an arrow;

FIG. 9 is an image illustrating a rat cornea implanted with lung micro-organs from a syngeneic rat. The implanted micro-organ (marked with arrow) is surrounded by newly formed blood vessels;

15 FIGs. 10A-B schematically illustrate a device for micro-organ preparation and delivery, in accordance with a preferred embodiment of the present invention;

FIG. 11 schematically illustrates a tissue scraper, in accordance with a preferred embodiment of the present invention;

20 FIG. 12 schematically illustrates the tissue scraper, in accordance with a preferred embodiment of the present invention;

FIGs. 13A-B schematically illustrate a tissue cutter, in accordance with a preferred embodiment of the present invention;

FIGs. 14A-B schematically illustrate the tissue cutter, in accordance with a preferred embodiment of the present invention;

FIG. 15 schematically illustrates the tissue cutter, when cutting is complete, in accordance with a preferred embodiment of the present  
5 invention;

FIGs. 16A-B schematically illustrate applying a medium for keeping micro-organs moist, in accordance with a preferred embodiment of the present invention;

FIGs. 17A-E, schematically illustrate the steps in inserting micro-  
10 organs into implanting devices, in accordance with a preferred embodiment of the present invention;

FIGs. 18A-C schematically illustrate the steps in implanting the micro-organs in a body, in accordance with a preferred embodiment of the present invention;

15 FIGs. 19A-C illustrate angiogenesis in implanted skin micro-organs (SMOs) 1, 3 and 7 days following implantation (arrows indicate newly formed blood vessels);

FIGs. 20A-B illustrate Regional blood flow in implanted SMOs (Figure 20A) as compared to flow induced by lung MO (Figure 20B).  
20 Fluorescent beads were used to determine the flow intensity;

FIGs. 21A-B illustrate vessel formation in young vs. old SMOs one month following implantation in young mice;

FIGs. 22A-B illustrate blood flow in young vs. old SMOs two weeks following implantation in young mice;

FIGs. 23A-G are photographs taken under a fluorescent microscope illustrating vessel formation in muscle tissue devoid of implanted SMOs (Figures 23A, G) and SMO implanted muscle tissue (Figures 23B-D); and

FIG. 24 illustrates blood vessel formation in a single SMO rescued seven days following implantation in the recipient rabbit.

#### DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present invention is of a method, extract, and pharmaceutical composition for inducing angiogenesis in a tissue of a mammal, and a device for the preparation and delivery of micro-organs into a mammal.

The principles and operation of the present invention may be better understood with reference to the drawings and accompanying descriptions.

Before explaining at least one embodiment of the invention in detail, it is to be understood that the invention is not limited in its application to the details of construction and the arrangement of the components set forth in the following description or exemplified in the examples section that follows. The invention is capable of other embodiments or of being practiced or carried out in various ways. Also, it is to be understood that the phraseology and terminology employed herein is for the purpose of description and should not be regarded as limiting.

As used herein, the term "micro-organ" refers to organ tissue which is removed from a body and which is prepared, as is further described below, in a manner conducive for cell viability and function. Such preparation may include culturing outside the body for a predetermined time period. The term

5 "angiopump" refers to micro-organs processed, preferably verified for cell viability and prepared in a manner ready, but not necessarily utilized, for immediate administration.

Complex multicellular organisms rely on a vascular network to support the needs of each and every cell for oxygen, nutrients and waste removal.

10 This complex network of blood vessels is created and sustained through the process of angiogenesis. In humans, the deterioration of the vascular network leads to occlusive arterial disease, which is the leading cause for morbidity and mortality in the Western world. Most currently available therapeutic options are based on surgical or other invasive procedures, such as vascular

15 bypass or angioplasty. These solutions are for the most part successful but may be short lived or not applicable to all patients. Since angiogenesis is a fundamental component of tissue and organ genesis, most tissues retain the capacity to induce new vessel formation during regeneration. Thus, the inventors of the present invention postulate that tissue which is removed from

20 the body is in essence at least attempting to undergo regeneration and thus can be utilized as an angiogenic stimulant, or more broadly for stimulation of cell growth processes.

The present invention provides a new approach to induce angiogenesis and other cell growth properties, which approach is based on the use of micro-organs. Such micro-organs retain the basic micro-architecture of the tissues of origin while at the same time are prepared such that cells of an organ explant are not more than 100-450 micron away from a source of nutrients and gases. Such micro-organs function autonomously and remain viable for extended period of time both as ex-vivo cultures and in the implanted state. Furthermore such micro-organs not only function but secrete a whole repertoire of angiogenic factors which induce a significant vascular network in their vicinity.

It will be appreciated, that although micro-organs can be utilized immediately following preparation, in some cases culturing outside the body for extended periods of time may be advantageous in order to increase viability. For example, in cases where soluble molecules are to be extracted, culturing of micro-organs is performed for predetermined time periods, which can be as short as 4 hours or as long as days or weeks.

Thus, the use of these micro-organs or extracts derived therefrom for inducing angiogenesis and other cell growth properties is dependent on the preservation of cellular function for various periods of time, prior to implantation. The present invention is based, in part, upon the discovery that under defined circumstances, growth of cells in different tissue layers of an organ explant, e.g., mesenchymal and epithelial layers, can be activated to proliferate, differentiate and function in culture.

The cell-cell and cell-matrix interactions provided in the explant itself are sufficient to support cellular homeostasis, thereby sustaining the microarchitecture and function of the organ for prolonged periods of time. As used herein, the term "homeostasis" is defined as equilibrium between cell proliferation and cell loss.

The support of cellular homeostasis preserves, for example, the natural cell-cell and cell-matrix interactions occurring in the source organ. Thus, orientation of the cells with respect to each other or to another anchorage substrate, as well as the presence or absence of regulatory substances such as hormones, permits the appropriate maintenance of biochemical and biological activity of the source organ. Moreover, the micro-organ can be maintained in culture without significant necrosis for at least 48 days.

*Source of explants for the micro-organ:*

Examples of mammals from which the micro-organs can be isolated include humans and other primates, swine, such as wholly or partially inbred swine (e.g., miniature swine, and transgenic swine), rodents, etc. Examples of suitable organs include, but are not limited to, liver, lung, other gut derived organs, heart, spleen, kidney, skin and pancreas.

*The growth media:*

There are a large number of tissue culture media that exist for culturing cells from animals. Some of these are complex and some are simple. While it is expected that micro-organs may grow in complex media,

it has been shown in United States Patent Application Serial Number 08/482,364 that cultures can be maintained in a simple medium such as Dulbecco's Minimal Essential Media (DMEM). Furthermore, although the micro-organs may be cultured in a media containing sera or other biological  
5 extracts such as pituitary extract, it has been shown in United States Patent Application Serial Number 08/482,364 that neither sera nor any other biological extract is required. Moreover, the micro-organ cultures can be maintained in the absence of sera for extended periods of time. In preferred embodiments of the invention, growth factors are not included in the media  
10 during maintenance of the micro-organ cultures *in vitro*.

The point regarding growth in minimal media is important. At the present, most media or systems for prolonged growth of mammalian cells incorporate undefined proteins or use feeder cells to provide proteins necessary to sustain such growth. Because the presence of such undefined  
15 proteins can interfere with the intended end use of the micro-organs, it will generally be desirable to culture the explants under conditions to minimize the presence of undefined proteins.

As used herein the language "minimal medium" refers to a chemically defined medium, which includes only the nutrients that are required by the  
20 cells to survive and proliferate in culture. Typically, minimal medium is free of biological extracts, e.g., growth factors, serum, pituitary extract, or other substances, which are not necessary to support the survival and proliferation of a cell population in culture. For example, minimal medium generally

includes at least one amino acid, at least one vitamin, at least one salt, at least one antibiotic, at least one indicator, e.g., phenol red, used to determine hydrogen ion concentration, glucose, and at least one antibiotic, and other miscellaneous components necessary for the survival and proliferation of the  
5 cells. Minimal medium is serum-free. A variety of minimal media are commercially available from Gibco BRL, Gaithersburg, MD, as minimal essential media.

However, while growth factors and regulatory factors need not be added to the media, the addition of such factors, or the inoculation of other  
10 specialized cells may be used to enhance, alter or modulate proliferation and cell maturation in the cultures. The growth and activity of cells in culture can be affected by a variety of growth factors such as insulin, growth hormone, somatomedins, colony stimulating factors, erythropoietin, epidermal growth factor, hepatic erythropoietic factor (hepatopoietin), and other cell growth  
15 factors such as prostaglandins, interleukins, and naturally-occurring negative growth factors, fibroblast growth factors, and members of the transforming growth factor-beta family.

#### *Culture Vessel:*

The micro-organs may be maintained in any suitable culture vessel  
20 and may be maintained at 37 °C in 5 % CO<sub>2</sub>. The cultures may be shaken for improved aeration.

With respect to the culture vessel in/on, which the micro-organs are preferably provided, it is noted that in a preferred embodiment such a vessel



may generally be of any material and/or shape. A number of different materials may be used to form the vessel, including but not limited to: nylon (polyamides), dacron (polyesters), polystyrene, polypropylene, polyacrylates, polyvinyl compounds (e.g., polyvinylchloride), polycarbonate (PVC),  
5 polytetrafluorethylene (PTFE; teflon), thermanox (TPX), nitrocellulose, cotton, polyglycolic acid (PGA), cat gut sutures, cellulose, gelatin, dextran, etc. Any of these materials may be woven into a mesh.

Where the cultures are to be maintained for long periods of time or cryopreserved, non-degradable materials such as nylon, dacron, polystyrene,  
10 polycarbonate, polyacrylates, polyvinyls, teflons, cotton or the like may be preferred. A convenient nylon mesh which could be used in accordance with the invention is Nitex, a nylon filtration mesh having an average pore size of 210  $\mu\text{m}$  and an average nylon fiber diameter of 90  $\mu\text{m}$  (Tetko, Inc., N. Y.).

#### *Dimensions of the Explant:*

15 In addition to isolating an explant which retains the cell-cell, cell-matrix and cell-stroma architecture of the originating tissue, the dimensions of the explant are crucial to the viability of the cells therein, e.g., where the micro-organ is intended to be sustained for prolonged periods of time, such as 7-21 days or longer.

20 Accordingly, the dimensions of the tissue or organ are selected to provide diffusion of adequate nutrients and gases such as oxygen to every cell in the three dimensional micro-organ, as well as diffusion of cellular waste out of the explant so as to minimize cellular toxicity and concomitant death

due to localization of the waste in the micro-organ. Accordingly, the size of the explant is determined by the requirement for a minimum level of accessibility to each cell in the absence of specialized delivery structures or synthetic substrates. It has been discovered, as described in United States Patent Application Number 08/482,364 that this accessibility can be maintained if the surface to volume index falls within a certain range.

This selected range of surface area to volume index provides the cells access to nutrients and to avenues of waste disposal by diffusion in a manner similar to cells in a monolayer. This level of accessibility can be attained and maintained if the surface area to volume index, defined herein, as "Aleph or Aleph index" is at least about  $2.6 \text{ mm}^{-1}$ . The third dimension has been ignored in determining the surface area to volume index because variation in the third dimension causes ratiometric variation in both volume and surface area. However, when determining Aleph, a and x should be defined as the two smallest dimensions of the tissue fragment.

As used herein, "Aleph" refers to a surface area to volume index given by a formula  $1/x + 1/a$ , wherein  $x$  = tissue thickness and  $a$  = width of tissue in mm. In preferred embodiments, the Aleph of an explant is in the range of from about  $2.7 \text{ mm}^{-1}$  to about  $25 \text{ mm}^{-1}$ , more preferably in the range of from about  $2.7 \text{ mm}^{-1}$  to about  $15 \text{ mm}^{-1}$ , and even more preferably in the range of from about  $2.7 \text{ mm}^{-1}$  to about  $10 \text{ mm}^{-1}$ .

Examples of Aleph are provided in Table 1 wherein, for example, a tissue having a thickness (x) of 0.1 mm and a width (a) of 1 mm would have an Aleph index of 11 mm<sup>-1</sup>.

5

**TABLE 1**  
***Different values for the surface area to volume ratio index "Aleph", as a function of a (width) and x (thickness) in mm<sup>-1</sup>***

x (mm)	Values of Aleph				
	a = 1	a = 2	a = 3	a = 4	a = 5
0.1	11	10.51	10.33	10.2	10.2
0.2	6	5.5	5.33	5.25	5.2
0.3	4.3	3.83	3.67	3.58	3.53
0.4	3.5	3	2.83	2.75	2.7
0.5	3	2.5	2.33	2.25	2.2
0.6	2.66	2.16	2	1.91	1.87
0.7	2.4	1.92	1.76	1.68	1.63
0.8	2.25	1.75	1.58	1.5	1.45
0.9	2.11	1.61	1.44	1.36	1.31
1.0	2	1.5	1.33	1.25	1.2
1.2	1.83	1.3	1.16	1.08	1.03
1.3	1.77	1.26	1.1	1.02	0.96
1.6	1.625	1.13	0.96	0.88	0.83
2.0	1.5	1	0.83	0.75	0.7

10

Thus, for example, cells positioned deepest within an individual micro-organ are at least 80 microns, and not more than 375 microns, away from a nearest surface of the individual micro-organ. These measurements facilitate the preservation of *in vivo* architecture, while concurrently ensuring that no cell is farther than 225-300 microns from a source of gases and nutrients.

15

Without being bound by any particular theory, a number of factors provided by the three-dimensional culture system may contribute to its success.

First, the appropriate choice of the explant size, e.g., by use of the  
5 above Aleph calculations, provides appropriate surface area to volume ratio for adequate diffusion of nutrients to all cells of the explant, and adequate diffusion of cellular waste away from all cells in the explant.

Second, because of the three-dimensionality of the explant, various cells continue to actively grow, in contrast to cells in monolayer cultures,  
10 which grow to confluence, exhibit contact inhibition, and cease to grow and divide. The elaboration of growth and regulatory factors by replicating cells of the explant may be partially responsible for stimulating proliferation and regulating differentiation of cells in culture, e.g., even for the micro-organ, which is static in terms of overall volume.

15 Third, the three-dimensional matrix of the explant retains a spatial distribution of cellular elements, which closely approximates that found in the counterpart organ *in vivo*.

Fourth, the cell-cell and cell-matrix interactions may allow the establishment of localized microenvironments conducive to cellular  
20 maturation. It has been recognized that maintenance of a differentiated cellular phenotype requires not only growth/differentiation factors but also the appropriate cellular interactions.

While reducing the present invention to practice, and as is further described in the Examples section hereinbelow, it was discovered that when micro-organs are implanted in a recipient, they provide a sustained dosage of a complex repertoire of angiogenic and other growth factors and cytokines, thus leading to the formation of new blood vessels in the implanted tissues of the host. It was also discovered that micro-organs could reverse ischemia in host tissues in both normal and aging animals. In addition, it was also revealed that micro-organs cultured *in vitro* also express the same repertoire of angiogenic and other growth factors and cytokines.

Thus, according to one aspect of the present invention there is provided a method of inducing angiogenesis and cell growth in a tissue of a mammal, such as, for example a human being. The method is effected by implanting at least one micro-organ within the tissue of the mammal. Examples of tissue suitable for micro-organ implantation include but are not limited to, organ tissue or muscle tissue.

Such implantation can be effected via standard surgical techniques or via implanting of micro-organ preparations into the intended tissue regions of the mammal utilizing specially adapted syringes employing a needle of a gauge suitable for the administration of micro-organs.

The micro-organs utilized for implantation are preferably prepared from an organ tissue of the implanted mammal or a syngeneic mammal, although xenogeneic tissue can also be utilized for the preparation of the micro-organs providing measures are taken prior to, or during implantation,

so as to avoid graft rejection and/or graft versus host disease (GVHD). Numerous methods for preventing or alleviating graft rejection or GVHD are known in the art and as such no further detail is given herein.

It will be appreciated that to facilitate transplantation of the explants  
5 which may be subject to immunological attack by the host, e.g., where xenogenic grafting is used, such as swine-human transplantations, the micro-organ can be inserted into or encapsulated by rechargeable or biodegradable devices and then transplanted into the recipient subject. Gene products produced by such cells can then be delivered via, for example, polymeric  
10 devices designed for the controlled delivery compounds, e.g., drugs, including proteinaceous biopharmaceuticals. A variety of biocompatible polymers (including hydrogels), including both biodegradable and non-degradable polymers, can be used to form an implant for the sustained release of a gene product of the cell populations of the invention at a particular target site. The  
15 generation of such implants is generally known in the art. See, for example, Concise Encyclopedia of Medical & Dental Materials, ed. By David Williams (MIT Press: Cambridge, MA, 1990); the Sabel et al. U.S. Patent No. 4,883,666; Aebischer et al. U.S. Patent No. 4,892,538; Aebischer et al. U.S. Patent No. 5,106,627; Lim U.S. Patent No. 4,391,909; and Sefton U.S. Patent  
20 No. 4,353,888.

According to one preferred embodiment of the present invention, at least a portion of cells of the micro-organ includes at least one exogenous polynucleotide sequence. Such polynucleotide sequence(s) are preferably

stably integrated into the genome of these cells although transient polynucleotide sequences can also be utilized. It will be appreciated that such exogenous polynucleotides can be introduced into the cells of the micro-organ following explantation from the organ tissue of the mammal or alternatively the mammal can be transformed with the exogenous polynucleotides prior to preparation of organ tissue or organs. Methods for transforming mammalian cells are described in detail hereinbelow.

Such exogenous polynucleotide(s) can serve for enhancing angiogenesis or cell growth by, for example, up-regulating or down-regulating the expression of one or more endogenous angiogenic or growth factors or cytokines expressed within these cells. In this case, the polynucleotide(s) can include trans-, or cis-acting enhancer or suppresser elements which regulate either the transcription or translation of the endogenous angiogenic and/or growth factors or cytokines expressed within these cells. Numerous examples of suitable translational or transcriptional regulatory elements, which can be utilized in mammalian cells, are known in the art.

For example, transcriptional regulatory elements are cis or trans acting elements, which are necessary for activation of transcription from specific promoters (Carey *et al.* (1989), J. Mol. Biol., 209:423-432; Cress *et al.* (1991) Science, 251:87-90; and Sadowski *et al.* (1988), Nature, 335:563-564).

Translational activators are exemplified by the cauliflower mosaic virus translational activator (TAV). See, for example Futterer and Hohn (1991) EMBO J. 10:3887-3896. In this system a di-cistronic mRNA is produced. That is, two coding regions are transcribed in the same mRNA  
5 from the same promoter. In the absence of TAV, only the first cistron is translated by the ribosomes. However, in cells expressing TAV, both cistrons are translated.

The polynucleotide sequence of cis acting regulatory elements can be introduced into cells of micro-organs via commonly practiced gene knock-in  
10 techniques. For a review of gene knock-in/out methodology see, for example, United States Patent Nos. 5,487,992, 5,464,764, 5,387,742, 5,360,735, 5,347,075, 5,298,422, 5,288,846, 5,221,778, 5,175,385, 5,175,384, 5,175,383, 4,736,866 as well as Burke and Olson, Methods in Enzymology, 194:251-270, 1991; Capecchi, Science 244:1288-1292, 1989; Davies *et al.*, Nucleic  
15 Acids Research, 20 (11) 2693-2698, 1992; Dickinson *et al.*, Human Molecular Genetics, 2(8):1299-1302, 1993; Duff and Lincoln, "Insertion of a pathogenic mutation into a yeast artificial chromosome containing the human APP gene and expression in ES cells", Research Advances in Alzheimer's Disease and Related Disorders, 1995; Huxley *et al.*, Genomics, 9:742-750  
20 1991; Jakobovits *et al.*, Nature, 362:255-261 1993; Lamb *et al.*, Nature Genetics, 5: 22-29, 1993; Pearson and Choi, Proc. Natl. Acad. Sci. USA, 1993, 90:10578-82; Rothstein, Methods in Enzymology, 194:281-301, 1991; Schedl *et al.*, Nature, 362: 258-261, 1993; Strauss *et al.*, Science, 259:1904-



1907, 1993, WO 94/23049, WO93/14200, WO 94/06908 and WO 94/28123 also provide information.

Down-regulation of endogenous angiogenic and/or growth factors or cytokines can also be achieved via antisense RNA. In this case the exogenous  
5 polynucleotide(s) can encode sequences which are complementary to the mRNA sequences of the angiogenic and/or growth factors or cytokines transcribed in the cells of the micro-organ. Down regulation can also be effected via gene knock-out techniques.

Up-regulation can also be achieved by overexpressing or by providing  
10 a high copy number of one or more angiogenic and/or growth factor or cytokine coding sequences. In this case, the exogenous polynucleotide sequences can encode one or more angiogenic or growth factors or cytokines such as but not limited to VEGF, bFGF, Ang1 or Ang2 which can be placed under the transcriptional control of a suitable promoter of a mammalian  
15 expression vector. Suitable mammalian expression vectors include, but are not limited to, pcDNA3, pcDNA3.1(+/-), pZeoSV2(+/-), pSecTag2, pDisplay, pEF/myc/cyto, pCMV/myc/cyto, pCR3.1, and their derivatives, which are available from Invitrogen, pCI which is available from Promega, pBK-RSV and pBK-CMV which are available from Stratagene, pTRES which is  
20 available from Clontech.

Numerous methods are known in the art for introducing exogenous polynucleotide sequences into mammalian cells. Such methods include, but are not limited to, direct DNA uptake techniques, and virus or liposome

mediated transformation (for further detail see, for example, "Methods in Enzymology" Vol. 1-317, Academic Press). Micro-organ bombardment with nucleic acid coated particles is also envisaged.

It will be appreciated that the angiogenic and/or growth factors or  
5 cytokines expressed in micro-organs can be extracted therefrom as a crude or refined extract in a soluble phase and utilized directly, or as part of a pharmaceutical composition for local administration into host tissues, e.g., in order to induce angiogenesis or other cell growth processes. It will further be appreciated that since micro-organs express different levels of the various  
10 angiogenic and/or growth factors and cytokines at different time points following implantation or during culturing, one can extract soluble molecules from different micro-organ cultures at different time points, which when locally administered in a series, mimic the temporal expression of an implanted or cultured micro-organ.

15 Thus, according to another aspect of the present invention, there is provided another method of inducing angiogenesis or other cell processes in a tissue of a mammal. This method is effected by extracting soluble molecules from micro-organs and locally administering at least one predetermined dose of the soluble molecules extracted into the tissue of the mammal. Numerous  
20 methods of administering are known in the art. Detailed description of some of these methods is given hereinbelow with regards to pharmaceutical compositions.

As mentioned above and according to another preferred embodiment of the present invention the soluble extracts are included in a pharmaceutical composition which also includes a pharmaceutically acceptable carrier which serves for stabilizing and/or enhancing the accessibility or targeting of the  
5 soluble extract to target body tissues.

Examples of a pharmaceutically acceptable carrier include but are not limited to, a physiological solution, a viral capsid carrier, a liposome carrier, a micelle carrier, a complex cationic reagent carrier, a polycathion carrier such as poly-lysine and a cellular carrier.

10 The soluble extract, which constitutes the "active ingredient" of the pharmaceutical composition, can be administered to the individual via various administration modes.

Suitable routes of administration may, for example, include transmucosal or parenteral delivery, including intramuscular, subcutaneous  
15 and intramedullary implanting as well as intrathecal, direct intraventricular, intravenous, intraperitoneal, intranasal, and/or intraocular implanting.

Preferably, the composition or extract is administered in a local rather than a systemic manner, for example, via implanting directly into an ischemic tissue region of the individual.

20 Pharmaceutical compositions of the present invention may be manufactured by processes well known in the art, e.g., by means of conventional mixing, dissolving, granulating, dragee-making, levigating, emulsifying, encapsulating, entrapping or lyophilizing processes.

Pharmaceutical compositions for use in accordance with the present invention thus may be formulated in conventional manner using one or more physiologically acceptable carriers comprising excipients and auxiliaries, which facilitate processing of the active ingredients into preparations which, 5 can be used pharmaceutically. Proper formulation is dependent upon the route of administration chosen.

For implanting, the active ingredient may be formulated in aqueous solutions, preferably in physiologically compatible buffers such as Hank's solution, Ringer's solution, or physiological salt buffer. For transmucosal 10 administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art.

The composition described herein may be formulated for parenteral administration, e.g., by bolus implanting or continuous infusion. Formulations for implanting may be presented in unit dosage form, e.g., in 15 ampoules or in multidose containers with optionally, an added preservative. The compositions may be suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents.

Pharmaceutical compositions for parenteral administration include 20 aqueous solutions of the active ingredient in water-soluble form. Additionally, suspensions of the active ingredients may be prepared as appropriate oily or water based implanting suspensions. Suitable lipophilic solvents or vehicles include fatty oils such as sesame oil, or synthetic fatty

acids esters such as ethyl oleate, triglycerides or liposomes. Aqueous  
implanting suspensions may contain substances, which increase the viscosity  
of the suspension, such as sodium carboxymethyl cellulose, sorbitol or  
dextran. Optionally, the suspension may also contain suitable stabilizers or  
5 agents who increase the solubility of the active ingredients to allow for the  
preparation of highly concentrated solutions.

In addition, the composition of the present invention may be delivered  
via localized pumps, or time release reservoirs which can be implanted within  
ischemic tissues of the individual.

10 Since angiogenic and other growth factors and cytokines are typically  
secreted from producing cells, micro-organs can also be cultured in suitable  
media and the conditioned media which includes the secreted angiogenic  
factors can be collected at predetermined time points and utilized as described  
hereinabove with respect to the soluble extract.

15 Thus, according to yet another aspect of the present invention there is  
provided a method of inducing angiogenesis or other cell processes in a tissue  
of a first mammal. The method according to this aspect of the present  
invention is effected by culturing at least one micro-organ in a growth  
medium to thereby generate a conditioned medium, collecting the conditioned  
20 medium following at least one predetermined time period of culturing and  
administering at least one predetermined dose of the conditioned medium into  
the tissue of the first mammal to thereby induce angiogenesis or other cell  
growth processes in the tissue.

Preferably, the growth medium is a minimal essential medium (described hereinabove) which does not contain undefined proteins or other growth factors which may interfere with the intended function of the conditioned media or which may cause undesired reactions in the administered mammal.

It will be appreciated that the collected conditioned media can be processed using chromatographic techniques, such as affinity columns and the like, so as to yield a substantially pure preparations which include an array of angiogenic or other growth factors suitable for inducing angiogenesis or other cell growth processes when administered to a mammal.

It will further be appreciated that the conditioned medium and the soluble extract described herein can also be derived from micro-organs which include exogenous polynucleotides as described hereinabove. In such cases, if the exogenous polynucleotides utilized encode angiogenic or other growth factors or cytokines, the sequence of such exogenous polynucleotides is selected suitable for the intended administered mammal. For example, in cases where the soluble extract or conditioned medium is administered to human recipients, human or humanized exogenous polynucleotides are preferably utilized.

The micro-organs according to the teachings of the present invention can be utilized following preparation, or alternatively they can be cryopreserved and stored at  $-160^{\circ}\text{C}$  until use. For example, micro-organs can

be cryopreserved by gradual freezing in the presence of 10% DMSO (Dimethyl Sulfoxide) and 20% serum.

This can be effected, for example, by encapsulating the micro-organs within planar sheets, (e.g., a semi-permeable matrix such as alginate) and inserting these encapsulated micro-organs into a sealable sterile synthetic plastic bag of dimensions closely similar to that of the encapsulated micro-organs. The bag would contain one plastic tubing input at one end and one plastic tubing output at the opposite end of the bag. The sealed plastic bag containing the planar sheet with the micro-organs could then be perfused with standard culture medium such as Ham's F12 with 10% DMSO and 20% serum and gradually frozen and stored at  $-160^{\circ}\text{C}$ .

An important goal in cardiovascular medicine would be to replace surgical bypasses with therapeutic angiogenesis. Yet, in spite of the considerable efficacy observed when angiogenic factors were used in animal models of coronary or limb ischemia, the clinical results have been disappointing. Recently, it has been suggested that clinical failure may be due to the application of the angiogenic factor or the combination of factors utilized. The angiogenesis method of the present invention overcomes such limitations of prior art methods.

The present invention brings forth a novel approach, which recognizes that angiogenesis and other cell growth processes are complex, highly regulated and sustained processes, mediated by several regulatory factors. The results presented by the present invention provide a model, which allows

studying the induction of angiogenesis, and cell growth both in and out of the body, and, as such, allows for the establishment of a pattern of expression of key regulatory factors. The results presented herein show that implanted micro-organs express several key angiogenic and other cell growth factors in a coordinated manner, both in and out of the body. Furthermore, as shown by *in vivo* experiments, micro-organs function as genuine *angiopumps* not only by transcribing angiogenic and other growth factors, but also by inducing the formation of new blood vessels. Furthermore, the magnitude of the induction is such that the vessels formed are sufficient to irrigate the surrounding area and rescue artificially induced hypoxic tissue regions in mice and rats.

The model for ischemia in rats presented hereinbelow in the example section appears to mimic chronic ischemia since no irreversible damage has occurred. In untreated animals, the ischemia was apparent only after exertion. Presumably, there is enough collateral circulation to keep the limb viable but not enough to allow normal function when faced with an additional challenge. The implantation of micro-organs appears to have reversed this condition by increasing blood supply to ischemic regions. The results show a significant difference between the micro-organ-treated and the control groups which difference is undoubtedly due to the induction of angiogenesis and other cell growth processes by the micro-organs.

In the series of mouse *in vivo* rescue experiments presented herein the ischemic insult was increased. Mice have inferior collateral circulation to the hindlimbs due to less developed tail arteries as compared to rats. In this



group, signs of acute irreversible ischemic damage such as gangrene and auto-amputation, were detected in the control group. This finding suggests that the present invention may also be useful for salvage procedures, though further testing is warranted.

5 In an additional series of trials presented below the ischemic challenge was further increased by inducing ischemia in previously diseased animals. Again, irreversible ischemic damage occurred only in the control animals. The damage to the control animals was so severe that stress tests were deemed superlative. Though the sample size was small the differences were  
10 marked. These results are particularly important since they illustrate that micro-organs are capable of inducing angiogenesis and other cell growth processes even in tissues affected by some types of peripheral vascular disease.

Thus, the present invention provides methods and compositions for  
15 inducing and maintaining blood vessel formation and other cellular processes within host tissues for the purposes of stimulating cell growth, rescuing ischemic tissues and/or generating natural bypasses around blocked blood vessels.

The present invention provides methods and compositions for the  
20 development and production of viable, sterile *angiopumps* that can be administered quickly and easily in an outpatient setting. It will be appreciated that the procurement, testing and administration of the *angiopumps* can thus

be accomplished most easily, or alternatively, can be similarly stored for administration at a later stage.

A novel device for the preparation and delivery of micro-organs is further provided and disclosed by the present invention. A detailed disclosure  
5 of the device is provided under Example 7 of the Examples section that follows.

Additional objects, advantages, and novel features of the present invention will become apparent to one ordinarily skilled in the art upon examination of the following examples, which are not intended to be limiting.  
10 Additionally, each of the various embodiments and aspects of the present invention as delineated hereinabove and as claimed in the claims section below finds experimental support in the following examples.

### EXAMPLES

15 Reference is now made to the following examples, which together with the above descriptions, illustrate the invention in a non-limiting fashion.

Generally, the nomenclature used herein and the laboratory procedures utilized in the present invention include molecular, biochemical, microbiological and recombinant DNA techniques. Such techniques are  
20 thoroughly explained in the literature. See, for example, "Molecular Cloning: A laboratory Manual" Sambrook et al., (1989); "Current Protocols in Molecular Biology" Volumes I-III Ausubel, R. M., ed. (1994); Ausubel et al., "Current Protocols in Molecular Biology", John Wiley and Sons, Baltimore,

Maryland (1989); Perbal, "A Practical Guide to Molecular Cloning", John Wiley & Sons, New York (1988); Watson et al., "Recombinant DNA", Scientific American Books, New York; Birren et al. (eds) "Genome Analysis: A Laboratory Manual Series", Vols. 1-4, Cold Spring Harbor Laboratory Press, New York (1998); methodologies as set forth in U.S. Pat. Nos. 4,666,828; 4,683,202; 4,801,531; 5,192,659 and 5,272,057; "Cell Biology: A Laboratory Handbook", Volumes I-III Cellis, J. E., ed. (1994); "Current Protocols in Immunology" Volumes I-III Coligan J. E., ed. (1994); Stites et al. (eds), "Basic and Clinical Immunology" (8<sup>th</sup> Edition), Appleton & Lange, Norwalk, CT (1994); Mishell and Shiigi (eds), "Selected Methods in Cellular Immunology", W. H. Freeman and Co., New York (1980); available immunoassays are extensively described in the patent and scientific literature, see, for example, U.S. Pat. Nos. 3,791,932; 3,839,153; 3,850,752; 3,850,578; 3,853,987; 3,867,517; 3,879,262; 3,901,654; 3,935,074; 3,984,533; 3,996,345; 4,034,074; 4,098,876; 4,879,219; 5,011,771 and 5,281,521; "Oligonucleotide Synthesis" Gait, M. J., ed. (1984); "Nucleic Acid Hybridization" Hames, B. D., and Higgins S. J., eds. (1985); "Transcription and Translation" Hames, B. D., and Higgins S. J., eds. (1984); "Animal Cell Culture" Freshney, R. I., ed. (1986); "Immobilized Cells and Enzymes" IRL Press, (1986); "A Practical Guide to Molecular Cloning" Perbal, B., (1984) and "Methods in Enzymology" Vol. 1-317, Academic Press; "PCR Protocols: A Guide To Methods And Applications", Academic Press, San Diego, CA (1990); Marshak et al., "Strategies for Protein Purification and

Characterization - A Laboratory Course Manual" CSHL Press (1996); all of which are incorporated by reference as if fully set forth herein. Other general references are provided throughout this document. The procedures therein are believed to be well known in the art and are provided for the convenience of the reader. All the information contained therein is incorporated herein by reference.

### ***EXAMPLE 1***

#### ***Micro Organs***

##### ***Materials and Experimental Methods***

Approval for animal experiments was obtained from the Animal Care and Use Committee of the Faculty of Science of the Hebrew University.

##### ***Micro-organ preparation:***

Adult animals (C57Bl/6 mice or Sprague Dawley rats) were sacrificed by asphyxiation with CO<sub>2</sub> and the lungs were removed under sterile conditions. The lungs were kept on ice and rinsed once with Ringer solution or DMEM including 4.5 gm/l D-glucose. micro-organs were prepared by chopping the lungs with a Sorvall tissue chopper into pieces approximately 300  $\mu$ m in width. micro-organs were rinsed twice with DMEM containing 500 units/ml Penicillin, 0.5 mg/ml Streptomycin and 2 mM L-Glutamine (Biological Industries) and kept on ice until use.

***Micro-organ implantation:***

Adult C57Bl/6 mice were anesthetized using 0.6 mg Sodium Pentobarbitol per gram body weight. The mice were shaved, and an incision about 2 cm long was made in the skin at an area above the stomach. A hemostat was used to create subcutaneous "pockets" on both sides of the incision, and 8-9 micro-organs were implanted in each pocket; implantation was done by simply layering the micro-organs over the muscle layer. The incision was sutured and the animals were kept in a warm, lit room for several hours following which they were transferred to the animal house. Four animals were sacrificed at a time interval of either 4 hours, 24 hours, 72 hours or 7 days following implantation and the implanted micro-organs were dissected from surrounding tissues under a surgical microscope and utilized for RNA extraction. The extracted RNA was reverse transcribed and the resulting cDNA was used as a template for PCR analysis using standard methodology. The oligonucleotide primer sequences utilized in the PCR reaction, the expected product size and references are given in Table 2 .

**TABLE 2**  
***RT-PCR primer sequence and source***

Name	Genebank #	Sequence	Product
Ang2	AF004326.1	F: 5'-CGTGGGTGGAGGAGGGTGGAC-3' (SEQ ID NO:1) R: 5'-TGC GTCAAACCACCAGCCTCC-3' (SEQ ID NO:2)	400 bp
$\beta$ -Actin		F: 5'-TACCACAGGCATTGTGATGG-3' (SEQ ID NO:3) R: 5'-AATAGTGATGACCTGGCCGT-3' (SEQ ID NO:4)	310 bp ****
Ang1	U83509	F: 5'-GGTC' (SEQ ID NO:5)	273 bp*/**

		R: 5'-CCAAGGGCCGGATCAGCATGG-3' (SEQ ID NO:6)	
VEGF	U41383	F: 5'-ACTTTCTGCTCTCTTGGGT-3' (SEQ ID NO:7)	444, 573**/**
		R: 5'-CCGCCTTGGCTTGTCACA-3' (SEQ ID NO:8)	

\* Another discrete band is often detected at approximately 320 bp – the origin of this band is unknown. \*\* Primers (Ang1) or primer sequence (VEGF) was kindly supplied by professor Eli Keshet, Israel. \*\*\*VEGF mRNA undergoes alternative splicing. PCR product sizes are 444 bp for VEGF<sub>121</sub>, 573 bp for VEGF<sub>165</sub>, and 645 bp for VEGF<sub>189</sub>. \*\*\*\* Ibrahim et al. 1998 *Biochimica et Biophysica Acta* 1403, 254-264.

### *Densitometric analysis and quantification:*

A 10 µl aliquot of each PCR reaction was electrophoresed in a 1.5 % agarose gel stained with ethidium bromide. Gels were imaged utilizing a Macintosh Centris 660 AV computer and a Fujifilm Thermal Imaging System with a Toyo Optics TV zoom lens (75-125 mm, F=1.8, with a Colkin orange 02 filter). Densitometric analysis was performed using the public domain NIH 1.61 analysis software. Quantitation was done by normalizing the expression level of each PCR product to those obtained for β-Actin. All PCR reactions were performed in duplicate. Statistical analysis of the relative expression levels of the various VEGF isoforms was performed by comparison of the means, using Welsh's *t*-test on nonpaired samples before and after implantation.

### *Ischemic tissue rescue experiments:*

32 Sprague Dawley rats aged 1-4 months and weighing 200-300 grams were utilized. The left common Iliac of each rat was ligated and excised from rats anesthetized using 0.9-1.1 mg/gram body weight of Pentothal at the aortic bifurcation just proximal to the Iliac bifurcation.

Sixteen rats were implanted with 3-4 micro-organs each 24 hours following the induction of ischemia. The micro-organs were implanted intramuscularly and subcutaneously along the Femoral artery (medially) and along the sciatic nerve (laterally). The remaining sixteen rats underwent sham implantation 24  
5 hours following the induction of ischemia.

Twenty six C57Bl/6 mice aged 1-3 months and weighing 19 to 27 grams were also tested. The left Common Iliac artery of anesthetized mice was ligated and excised at the aortic bifurcation just proximal to the Iliac bifurcation. 3-4 micro-organs were implanted in each mouse at 24 hours  
10 following the induction of ischemia. Nine mice were implanted intramuscularly and subcutaneously along the Femoral artery (medially) and along the sciatic nerve (laterally) in the proximal left hindlimb. Seventeen control mice were prepared for implantation following ischemia induction but no implantation was performed. Animals that had venous or nervous damage  
15 during the operation as well as those that suffered from significant bleeding were excluded from the trial.

Seven old C57Bl/6 mice aged 22 months and weighing 24 to 28 grams also underwent ligation and excision of the left Common Iliac artery as described above. Three were immediately implanted with micro-organs  
20 derived from normal healthy syngeneic mice. Four had immediate sham implantation. None suffered from venous or nervous damage or had significant bleeding during the operation.

***Functional assay:***

The animals were tested on the first and second days following implantation to rule out nerve damage. The test consisted of swimming in a lukewarm water bath, which was set at a water level such that the animal  
5 needed to constantly exert all four limbs in order to stay afloat. The time limits for exercise were gradually increased. During the first week the time limit was 3 minutes or until efforts to remain afloat ceased. During the second week the limit was raised to 5 minutes, while from the third week onwards the time limit was 6 minutes. A scale from 0 to 10 was created to  
10 assess the degree of claudication. A score of 0-1 indicated normal or near normal gait. A score of 2-3 meant slight to moderate claudication with normal weight bearing. A score of 4-5 indicated moderate claudication with disturbance in weight bearing. A score of 6-7 indicated severe claudication.

A score of 8-9 indicated a non functioning limb, atrophy or contracture  
15 and a score of 10 meant gangrene or autoamputation. The scores were assigned by an independent observer not involved in the experiment and having no knowledge of previous animal treatments.

***Angiography:***

Angiography was performed on several rats at days 4, 14, 26 and 31  
20 following implantation. The rats were anesthetized as previously described and a P10 catheter was introduced through the right superficial femoral artery and placed in the aorta. A bolus implanting of 1 cc Telebrix was injected and



the animal was photographed every 0.5 seconds. Animals undergoing angiography were subsequently excluded from the trial groups.

### *Experimental Results*

#### *Implanted micro-organs induce angiogenesis:*

5        Figure 1 illustrates the response of surrounding tissues to implanted micro-organs. When a micro-organ is implanted subcutaneously into a syngeneic animal, it induces an angiogenic response towards the micro-organ (arrow, Figure 1). A major blood vessel forms and branches into smaller vessels, which branch into a net of capillaries, which surround the implanted  
10    micro-organ.

micro-organs transcribe a sustained and dynamic array of angiogenic growth factors when implanted subcutaneously into syngeneic mice. Figure 2 illustrate a representative semi-quantitative analysis of several known angiogenic growth factors as determined from the RT-PCR analysis  
15    performed on RNA extracted from the micro-organs. As seen from the results, a strong induction of angiogenic factor expression occurs at 4 hours post implantation (PI). Following this initial induction, each individual growth factor follows a different expression pattern as is further detailed below.

20        *VEGF*: VEGF transcription level continued to rise at 24 hours PI. At three days PI, transcription levels of VEGF decreased. In the days following, lower mRNA levels of this angiogenic factor were detected, which levels were probably necessary in order to maintain the neo-angiogenic state thus

formed. At seven days PI, VEGF mRNA returned to a level similar to that detected in micro-organs at the time of implantation ( $t_0$ ).

*Angiopoietin 1:* The level of Ang1 mRNA increased for the first 4 hours PI, although variation was high. At one to three days PI, transcription  
5 dropped to levels which are even lower than that detected for micro-organs at the time of implantation ( $t_0$ ) (see Maisonpierre et al., 1997, *Science* 277, 55-60, Gale and Yancopoulos, 1999 *Ibid.*). At seven days PI, Ang1 mRNA returned to a level similar to that detected at  $t_0$ .

*Angiopoietin 2:* Ang2, the antagonist of Ang1, was transcribed at  
10 high levels at 24 hours PI. mRNA levels dropped at 3 and 7 days PI, although these levels were still higher than the levels detected at  $t_0$ , possibly due to ongoing vascular remodeling in and around the implanted micro-organ.

Thus, as is evident from these results, implanted micro-organs transcribe a dynamic array of factors, both stimulators and inhibitors, which  
15 participate in the regulation of angiogenesis. This transcription pattern which is responsible for the generation of new blood vessels around the micro-organs is sustained over a period of at least one week PI.

*Micro-organs transcribe a sustained and dynamic array of angiogenic growth factors when cultured:*

20 In order, to determine the capacity of micro-organs to transcribe angiogenic factors when cultured ex-vivo, micro-organs prepared as described above, were grown in the absence of serum for periods of over one month. Samples were removed at various time points and assayed for the

mRNA levels of the several factors. Figure 4 illustrate a representative semi-quantitative analysis of several known angiogenic growth factors as determined from RT-PCR performed on RNA extracted from cultured micro-organs (Figure 3). As shown in both Figures a strong induction of angiogenic  
5 factor expression occurs 4 hours following culturing. Following this initial induction, each different growth factor follows a different expression pattern as is described in detail below.

**VEGF:** VEGF expression levels continued to rise 24 hours after culturing. 3 days after culturing, the expression level of VEGF decreased  
10 only to increase again at 7 days PI. In the following days expression levels drop and VEGF expression returns to a level comparable to that expressed by micro-organs at the time of culturing.

**Angiopoietin 1:** The level of Ang1 expression increased for the first 4 hours following culturing although variation was high. Expression dropped 1  
15 to 3 days after culturing to levels even lower than that detected at time of culturing. At seven days after culturing Ang1 expression returned to a level comparable to the level at time of culturing.

**Angiopoietin 2:** Ang2, the antagonist of Ang1, was expressed at a high level during the first day after culturing. The expression levels were  
20 lower 3 and 7 days after culturing, although they are still higher than the expression level at time of culturing.

As is evident from these results, micro-organs which are cultured outside the body remain viable and functional for over a month *in vitro* and

express a dynamic array of angiogenic factors, including both stimulators and inhibitors, which participate in the regulation of angiogenesis.

*Implantation of micro-organs reverse ischemia in limbs of rats and mice:*

5        *Series 1:* The left common iliac artery of thirty two rats was ligated and excised as described above. micro-organs implantation was conducted in sixteen of these rats while the sixteen remaining rats served as the control group (sham operations). All 32 rats survived the operation. No significant difference was detected between the two groups prior to exertion (Figure 5).  
10 Following exertion, a significant difference was detected; the cumulated average claudication score for the control group was 4.8 whereas in the micro-organ implanted group the score was 1.6 (Figure 6). Similar results were recorded throughout the study period. The control group scored 5 on days 6-10 post operation (PO), 5 at 11-15 days PO and 4 at day 17 PO. The  
15 scores for the micro-organ implanted group were 1.67, 1.5 and 1.7, respectively. It should be noted that the micro-organ implanted group included one rat with an average score of 6.5. A histological examination revealed necrotic micro-organ implants in this rat.

*Series 2:* Twenty six young C57Bl/6 mice were operated as described  
20 above without operative damage or preoperative mortality. micro-organ implantation was conducted in nine of these mice while seventeen served as control (sham operations). Of the 17 control mice, 4 developed gangrene on the ischemic-induced limb and died 2-3 days PO (23.5 %). Another mouse

from this group had autoamputation of an atrophied limb 8 days after operation (5.9 %). None of the micro-organ implanted mice developed gangrene, autoamputation or postoperative death (0 %). The average cumulated post exertion claudication score for the control group was 6 with  
5 scores of 7.7 on days 5-9 PO, 6.2 on days 13-19 PO and 4.1 on days 21-25 PO. The average cumulated claudication score for the micro-organ implanted group was 2.4, with scores of 1.8 on days 5-9 PO, 2.2 on days 13-19 PO and 3.1 on days 21-25 PO (Figure 7).

***Micro-organ implantation rescues ischemic limbs in old mice:***

10       **Series 3:** Seven aged C57Bl/6 mice were operated upon with no operative damage or death. Three mice received micro-organ implants and 4 served as control. Of the control group, 1 developed gangrene and died 3 days PO (25%) and one had autoamputation of an atrophied limb 5 days PO (25%). The remaining two mice had non functioning limbs at rest (a score of 8 on the  
15 claudication index). None of the micro-organ implanted mice developed gangrene or autoamputation (0%) and their average claudication score at one week was 5.7.

***Implanted micro-organs are viable, and vascularized:***

20       In sampled rat specimens the micro-organ implants were viable, with preserved architecture and no evidence of rejection. The micro-organs and surrounding muscle tissue was vascularized via macroscopically visible blood vessels.

*Angiography reveals angiogenic activity in micro-organ-implanted rats:*

Angiography was performed on days 4, 14, 26 and 31 PO. There were subtle but detectable differences between the micro-organ-treated groups and the control groups. Evidence of increased angiogenic activity in the implanted limb was detected as early as day 4 PO. New, medium sized blood vessels were visible in the implanted limb sixteen days PO.

*EXAMPLE 2*

10

*Spleen micro-organs*

Mouse Spleen micro-organs were prepared from as described hereinabove and implanted into syngeneic mice. Figure 8 illustrates a micro-organ (arrow) which was implanted subcutaneously into the syngeneic mouse and examined at six months following implantation. As is clearly demonstrated in Figure 8, the micro-organ induced angiogenesis. In fact, the pattern of blood vessels formed gives the impression that the micro-organ is micro-organ was an inherent organ of the host.

*EXAMPLE 3*

20

*Cornea implantation of micro-organs*

The cornea is the only tissue of the body, which is devoid of blood vessels. As such, the cornea is an excellent model tissue for studying angiogenesis. Rat lung micro-organs were implanted in the corneas of

syngeneic rats. As shown in Figure 9, a most remarkable angiogenic pattern was also induced in the cornea. These remarkable results again verify that micro-organs are effective in inducing and promoting angiogenesis.

5

#### **EXAMPLE 4**

##### ***Mouse Skin MOs implanted in C57BL mice***

##### ***Materials and Experimental Methods:***

Adult C57BL mice were anesthetized using Sodium Pentobarbital. The mice were shaved, and an incision about 1cm long was made in the skin at the center of the stomach. A haemostat was used to create subcutaneous  
10 pockets on both sides of the incision, and about 10 skin MOs (SMOs), prepared as described earlier, were placed side by side (on their side thus exposing all tissue layers) in each pocket. The incision was closed using surgical sutures. One, three, seven and thirty days following implantation, the  
15 recipient mice were sacrificed and the SMOs were excised from surrounding tissues under a surgical microscope (Figures 19A-C). Ten skin MOs were taken at time zero.

##### ***Determination of regional blood flow:***

SMO implanted Mice were anesthetized using Sodium Pentobarbital  
20 (0.06 mg per gram body weight) and the right carotid artery was cannulated using heparinized-saline (20 U/ml) filled PE-10 tubing which was narrowed at the portion inserted into the vessel. The tubing was utilized to inject  $10^5$  polystyrene yellow-green fluorescent microspheres (Molecular probes 15  $\mu$ m

in diameter) into the left ventricle and 0.15 ml saline which was slowly injected into the left ventricle over a period of 30 seconds following injection of the microspheres.

The microspheres were found distributed throughout the implanted skin micro-organ indicating that blood was flowing into the SMO and that the vascular network had further expanded throughout the whole tissue (Figure 20A).

### EXAMPLE 5

10

#### *An aging model*

It is a well known fact that as individuals age the risk of cardio and peripheral vascular diseases such as atherosclerotic increases while regenerative capabilities responsible for wound healing among other processes decrease.

15

One factor which may contribute to this increase in risk and decrease in regenerative capabilities is a decrease in the body's capacity to stimulate angiogenesis.

To verify this theory, SMOs from old mice were compared to SMOs from young mice as far as their capacity to stimulate angiogenesis in a recipient host.

20

#### *Materials and Experimental Methods:*

*RNA extraction and cDNA synthesis:* Total RNA was extracted from equal amount of skin MOs using the acid-guanidine-phenol method



described by Chomczynski, P. (1994) in *Cell biology: a laboratory handbook*, ed. E, C. J. (Academic press, Vol. 1, pp. 680-683 Chomczynski.) Additionally, cDNA was synthesized from 1-2 µg total RNA with poly-d(T)<sub>12-18</sub> primer, obtained, for example, from Promega USA, and Moloney murine leukemia virus reverse transcriptase, obtained for example, from Promega USA.

***Reverse transcription (RT) PCR analysis:***

1 µl cDNA samples were subjected to PCR amplification in 1.5 mM MgCl<sub>2</sub>. The number of PCR cycles was 36 for all angiogenic factors but actin which was amplified using 24 cycles. For each series of primers, a positive control PCR reaction using cDNA synthesized from lung MOs mRNA extracted at time zero and a negative reaction using no template were also performed. The same primers, as described hereinabove in Table 2, were used.

***Experimental Results:***

***Mouse Skin MOs (SMOs) implants in C57BL mice in vivo:*** The negative controls did not yield a detectable signal. Skin MOs transcribed all of the angiogenic factors tested (Ang1, Ang2, HGF, bFGF, three isoforms of VEGF, Ephrin 3b, and Mef2C) exhibiting expression kinetics somewhat different than that of lung MOs. In addition, angiogenic induction activity of skin MOs was at least as strong as that exhibited by lung MOs.

Comparison of angiogenesis between SMO made of old(2 years old) and young(2 months old) mice skin, one month following implantation in a

young mouse, revealed that no decrease in vessels formation can be detected in the old SMO (Figures 21A-B).

In addition, comparison of blood flow in SMOs made of old (2 years old) and young (2 months old) mice, 2 weeks following implantation in a young mouse, revealed that blood flow in the old SMOs implanted mouse was as high as, if not higher than, that of the young SMOs implanted mouse (Figures 22A-B).

Thus, it is clear that SMOs made of the old mouse did not lose the capability to induce angiogenesis.

10

#### **EXAMPLE 6**

##### ***Skin MOs implantation into Rabbits***

This study utilizes the methodology described hereinabove to stimulate angiogenesis in rabbits. Since a rabbit is a larger animal it can be used to more accurately model the process of angiogenesis induction in humans.

##### ***Materials and Experimental Methods:***

Rabbits weighing approximately 2.5 kg each were anesthetized and a piece of skin from the center the stomach was excised and used to prepare SMOs in a manner similar to that described above for mouse SMOs. Four SMOs were implanted 5-8 mm apart in a straight line within the muscle tissue of each of the rabbit legs. Seven days following implantation, a blood flow

distribution assay was performed on each rabbit using the microspheres and methodology described hereinabove.

***Experimental Results:***

As shown in Figures 23A-G, the injected microspheres were found  
5 distributed throughout the implanted SMOs indicating that blood was flowing  
into the SMOs and that the vascular network had further expanded throughout  
the whole tissue.

In addition, the average amount of beads found in unimplanted muscle  
tissue (Figures 23A and G) was much lower than that of SMO implanted  
10 muscle tissue (Figures 23B-F).

Following blood flow determination, a single SMO was removed and  
the regional blood flow reaching directly into the SMO was determined by  
measuring the fluorescence intensity of the SMO. Negative control non-  
viable SMOs were found to yield non-detectable fluorescence and no  
15 fluorescent beads were observed inside the dead SMOs. In contrast, as shown  
in Figure 24, a single viable SMO induced a significant amount of blood  
vessel formation as exemplified by the significant number of green  
fluorescent beads observed seven days following implantation into the  
recipient rabbit.

**EXAMPLE 7*****Device for the preparation and delivery of micro-organs***

The present invention relates also to a device for the preparation and delivery of micro-organs, such as angiopumps for inducing angiogenesis in a tissue of a mammal.

Referring further to the drawings, Figure 10A schematically illustrates a device 10 for micro-organ preparation and delivery, in accordance with a preferred embodiment of the present invention. Device 10 includes:

- i. a tissue scraper 20, for obtaining a tissue biopsy;
- 10 ii. a tissue cutter 40, for cutting the tissue biopsy into a plurality of fragments, forming micro-organs; and
- iii. at least one implanting device 60, arranged within an implanting chamber 70 and detachably coupled to device 10, for receiving a single micro-organ, when coupled to device 10, and for implanting the micro-
- 15 organ into a subject (not shown), after decoupling from device 10.

Additionally, device 10 includes a casing 22, a base 21A, and a ramp 21B, which together form an enclosure that may be sealed. Device 10 is preferably about 100 mm in width and about 300 mm in length. It will be appreciated that somewhat larger or smaller dimensions are also possible.

20 Referring further to the drawings, Figure 10B schematically illustrates a control system 12 for device 10. Control system 12 may be a PC computer, a laptop, a palm computer, or the like, or a dedicated control system for device 10, having a processor and preferably a memory. Preferably, control

system 12 includes a control panel 13, having several knobs or buttons 14, a keyboard 11, a display panel 15, which may include an interactive display panel, at least one light 16, for indicating that the system is on, one or more warning lights 17, for example, to indicate that the temperature has exceeded  
5 a recommended value, or that a travel mechanism is jammed, and a read and preferably write device 9, such as a diskette drive, a CD drive, a minidisk drive, or the like, for running or recording a predetermined sequence of tasks. Additionally, control system 12 may control a plurality of devices 10 at any one time. Communication between one or more devices 10 and control  
10 system 12 may be wired or wireless.

Alternatively, no control system is used, but some functions of device 10 are automated and controlled by knobs 38 on device 10 (Figure 10A). Alternatively or additionally, buttons or switches 38, or the like may be used on device 10.

15 Referring further to the drawings, Figures 11 and 12, together with Figure 10A, schematically illustrate tissue scraper 20, in accordance with a preferred embodiment of the present invention. Tissue scraper 20 may be for example, a standard, manually operated dermatome such as that manufactured by Robbins Instruments Inc. or by Aesculap<sup>®</sup> or a similar, preferably  
20 electrical dermatome.

Preferably, tissue scraper 20 includes a scraping blade 24 (Figure 12), adapted for scraping a split-thickness skin biopsy (SPS) 25. A split-thickness biopsy is usually obtained by cutting, for example with a commercially

available dermatome, parallel to the surface of the organ, a flat organ explant of predetermined thickness. The position of the blade determines the depth of the cutting and thus the thickness of the flat biopsy. Several illustrations of SPSs of varying thickness can be found in the literature (for example see  
5 Kondo S, Hozumi Y, Aso K., Long-term organ culture of rabbit skin: effect of EGF on epidermal structure in vitro. J Invest Dermatol. 1990 Oct;95(4):397-402.

Additionally, tissue scraper 20 includes casing 22 and ramp 21B. Casing 22 includes a movable portion 18, which may be raised and lowered,  
10 as shown by arrow 23. When raised, it exposes a window 19 (Figure 11), through which SPS 25 is admitted. Additionally, movable portion 18 includes a guillotine-like blade 26, which when lowered, cuts SPS 25 off the body. The movement of movable portion 18 may be manual, or may be controlled from control system 12, or by one of switches 38 of device 10.

15 Preferably, scraping blade 24 is adapted for cutting SPS 25 of a width A of preferably 6-8 mm (Figure 11). A length B of SPS 25 is approximately 2 cm (Figure 10A). A thickness C of SPS 25 may be about 1.0 - 1.4 mm, and preferably not less than about 650 microns (Figure 11). By selecting scraping blade 24 of a predetermined width A, and by lowering guillotine-like blade 26  
20 after a predetermined length has been scraped, both width A and length B may be predetermined. Additionally, by adjusting the a distance R between scraping blade 24 and ramp 21B, thickness C may be predetermined. In other words, the height of the blade 24 can be lowered or raised with respect to

21B, thus affecting the thickness of the SPS. It will be appreciated that scraping blade 24 may be replaceable, for example, with a blade generating a different width A. Alternatively or additionally, blade 24 may be replaced when it grows dull.

5 Preferably, a region of the body 27 (Figure 11) from which SPS 25 may be scrapped is the stomach of the patient. Alternatively, region 27 may be the back of the arm, the buttocks, the hips or another area, which is generally unexposed, and which is generally denuded of hairs. It will be appreciated that SPS 25 may be taken from another person, acting as a donor,  
10 rather than from the patient. Additionally, it will be appreciated that SPS 25 may be taken from mammals, such as primates, swines, such as wholly or partially inbred swines (e.g., miniature swines, and transgenic swines), rodents, and the like.

Prior to the scraping, region 27 is shaved, thoroughly cleaned, and  
15 disinfected using standard surgical procedures. Similarly, device 10 is thoroughly sterilized. In a preferred embodiment device 10 is disposed after use.

As seen in Figure 11, the scraping operation is manual. A hand 8 of an operator pushes device 10 into region 27 to scrape a tissue biopsy.

20 As seen in Figure 12, when movable portion 18 is lowered, and guillotine-like blade 26 cuts SPS 25 off, a sealed enclosure 30 is formed around SPS 25. At least two conveyer belts strips 28, arranged on rollers 29 (Figure 10A), transfer SPS 25 into sealed enclosure 30, without human

contact. It will be appreciated that other automated means of transferring SPS 25 may be employed, for example, a wide conveyer belt, whose width is wider than width A of SPS 25. Alternatively, a rigid platform, seated on a moving gantry, may be used. Alternatively, other automated means of  
5 transferring SPS 25 may be employed, as known. Preferably, the transfer of transfer SPS 25 into sealed enclosure 30 is controlled from control system 12. Alternatively, it is controlled by one of switches 38 of device 10. Alternatively, conveyer belt 28 may be manually controlled by a winding handle, or a similar mechanism.

10        Additionally, as seen in Figure 12, device 10 includes washing apparatus 31, comprising a washing-solution dispenser 32, an inlet 35 and a drain 34. Dispenser 32 sprays an appropriate washing solution 36 over SPS 25, for thoroughly rinsing it. Washing solution 36 may be, for example, a standard culture medium DMEM with 500 units/ml Penicillin, 0.5 mg/ml  
15 Streptomycin. Washing solution 36 is admitted to dispenser 32 via inlet 35, and drains away through drain 34. Thus, rinsing takes place from the top of SPS 25. It will be appreciated that other means for rinsing SPS 25 may be employed. For example, a plurality of sprinklers may be used to spray SPS 25. Alternatively, SPS 25 may be soaked in a bath of washing solution 36,  
20 for a predetermined time. Preferably, the rinsing of SPS 25 is controlled from control system 12. Alternatively, it is controlled by one of switches 38 of device 10. Alternatively, device 10 is manually filled with washing



solution 36, and the rinsing and drainage of washing solution 36 is powered by gravity.

As seen in Figure 10A, following the rinsing, at least one second conveyer belt 42, and preferably two second conveyer belts 42, operating on  
5 rollers 39, transfer SPS 25 to tissue cutter 40, preferably, aseptically and preferably, without human direct intervention. It will be appreciated that other automated means of transferring SPS 25 may be employed, as was noted hereinabove. Similarly, control system 12, one of switches 38, or a manual control may be used for the automated transfer of SPS 25 to tissue  
10 cutter 40.

Referring further to the drawings, Figures 13A - 13B, 14A - 14D, together with Figure 10A, schematically illustrate tissue cutter 40, in accordance with a preferred embodiment of the present invention.

As seen in Figures 10A and 13A, at tissue cutter 40, conveyer belts 42  
15 transfer SPS 25 to region 41, wherein SPS 25 is supported by a plurality of rods 54, arranged in a single line, and forming micro-organ guides 54. Micro-organ guides 54 are preferably formed of medical grade polycarbonate of internal diameter approximately 0.4 mm and length approximately 16 cm. The internal diameter of micro-organ guides 54 is approximately 0.4 mm and  
20 their length is approximately 15 - 16 cm. It will be appreciated that somewhat smaller or larger values are also possible.

As seen in Figures 14A - 14B, micro-organ guides 54 have a first section 56 of a circular cross section and a second section 58, which is

formed as a half circle, having a concave inner surface 62. It will be appreciated that both sections 56 and 58 may be solid or hollow. However, in accordance with a preferred embodiment of the present invention, section 56 is hollow and section 58 is solid. Additionally, micro-organ guides 54  
5 include a position marker 68, a notch 64, and a distal edge 59. The purpose of position marker 68 and notch 64 will be illustrated hereinbelow, in conjunction with Figures 17A - 17E. Region 41, which supports SPS 25, is formed of half-split rods 58, which provide a solid flat or preferably a concave support for the SPS before being cut into micro-organs.

10        Additionally, as seen in Figures 13A and 13B, tissue cutter 40 includes a plurality of parallel, surgical-grade blades 44, arranged on a moving gantry 46, which is manually manipulated by a handle 48. Handle 48 protrudes from casing 22 through a slit window 50 which permits the manual control of gantry 46 and may further define its maximum travel.  
15        Preferably, gantry 46 glides along a straight edge 45. In accordance with an alternative embodiment, of the present invention, the travel of gantry 46 may be automated, and controlled from control system 12, or by one of switches 38 of device 10.

      In accordance with a preferred embodiment of the present invention,  
20        blades 44 are arranged at an angle with respect to SPS 25, as seen in Figures 13B and 14B. Alternatively, they may be rotatable disc-blades, similar to rolling pizza cutters, operative to cut as they roll. Alternatively, wire cutters, similar to cheese or egg cutters, may be used.

Preferably, plurality of blades 44 are adapted to operate simultaneously, as a single ensemble, and touch SPS 25 at all points at the same time, thus avoiding moving, wrinkling, or folding SPS 25 during cutting.

5        Blades 44 may be powered manually or by a motor. Alternatively, blades 44 may be spring loaded, and operate in a guillotine-like fashion. In accordance with an embodiment of the present invention, gantry 46 and blades 44 may be removable and replaceable, so that different types of blades 44 may be used at different times.

10        In accordance with the present invention, a distance  $d$  between adjacent blades 44 (Figure 13B) is substantially equal to, or smaller than a diameter  $e$  of half rods 58, which form micro-organ guides 54 (Figure 14B). In consequence, as seen in Figure 14B, as blades 44 cut SPS 25 to a plurality of fragments 66, each fragment 66 (possibly, except edge fragments 53 and  
15        55) is supported by 62 of one of half rods 58. Preferably, 11 blades are used, to cut 12 fragments 66. However, it will be appreciated that other numbers may similarly be employed.

A key feature of the present invention is distance  $d$  between adjacent blades 44. It forms the width of fragments 66. That distance is between  
20        and 750 microns, and preferably 300 microns, so as to ensure that cells positioned deepest within fragment 66 are at least 80 microns and not more than about 375 microns away from a nearest surface of fragment 66. The

nearest surface may be one of surfaces 63 and 65. Thus fragments 66 are operative as a micro-organ 66, or micro-organs 66.

It will be appreciated that thickness  $C$  (Figure 11) may also be less than 750 microns (although as noted, it is more than 650 microns) thus cells  
5 positioned deepest within micro-organ 66 may be less than 375 microns away from two nearest surfaces.

In accordance with an embodiment of the present invention, gantry 46 and blades 44 may allow adjustments of distance  $d$  between adjacent blades 44, so long that distance  $d$  remains smaller than diameter  $e$  of micro-organs  
10 54. Alternatively or additionally, gantry 46 and micro-organs 54 may be removable and replaceable, with others, of different parameters  $d$  and  $e$ .

Edge fragments 53 and 55, whose widths are generally smaller than  $d$ , are generally discarded. However, one edge fragment may be used for a viability test, as will be described hereinbelow, in conjunction with Figures  
15 16A and 16B. As seen in Figure 13B, first section 56 of micro-organ 54 preferably acts as a stop that prevents SPS 25 from sliding along concave surface 62 of one of half rods 58, by the force of blades 44.

Alternatively or additionally, as seen in Figures 14C and 14D, SPS 25 may be held in place, for example, by side clamps 52, or similar devices, that  
20 may close on edge fragments 53 and 55, as shown by arrows 57, to prevent wrinkling or sliding that may be brought about by the force of blades 44. Side clamps 52 may extend the width of conveyer belts 42 (Figure 13A),

while gantry 46 and blades 44 may operate within the span of conveyer belts 42.

Referring further to the drawings, Figure 15 schematically illustrates blades 44 and handle 48, when cutting is complete, in accordance with a preferred embodiment of the present invention. Preferably, when cutting is complete, blades 44 are raised from a position 49, between micro-organs 66, to a position 49', above micro-organs 66, by raising handle 48 from its operating position 49 to locking position 49'.

It will be appreciated that device 10 may be further operative as a sealed treatment chamber, in particular, at zone 41 (Figure 10A). Treatment may be performed prior to cutting or after it. Treatment may include incubation at a specific temperature, wherein device 10 may further include a heater/cooler 67 and a thermostat 69. Alternatively or additionally, treatment may include treating SPS 25 with a special solution or hormone, which may be introduced via washing apparatus 31. Treatment may be controlled from control system 12, by one of switches 38, or manually.

Referring further to the drawings, Figures 16A and 16B schematically illustrates applying a medium 71 for keeping micro-organs 66 moist, or for supplying nutrients, in accordance with a preferred embodiment of the present invention. Medium 71 is applied via washing apparatus 31, which may be coupled to first sections 56 of micro-organ guides 54 (Figure 14B), which in this case are formed as hollow tubes. These lead to second sections 58, wherein micro-organs 66 are supported.

Additionally or alternatively, micro-organs 66 may be rinsed via washing apparatus 31, in a similar manner.

In accordance with the present invention, treatment may further include culturing, which may require at least an hour.

5        Additionally or alternatively, treatment may include transformation. Transformation may comprise introducing to at least a portion of the cells of micro-organs 66 at least one exogenous polynucleotide sequence preferably selected for regulating angiogenesis. The at least one exogenous polynucleotide sequence may be integrated into a genome of the portion of  
10   the cells of micro-organs 66.

The at least one exogenous polynucleotide sequence may be designed for regulating expression of, for example, at least one angiogenic factor of a plurality of angiogenic factors. Additionally, the at least one exogenous polynucleotide sequence may include an enhancer or a suppresser sequence.  
15   Furthermore an expression product of the at least one exogenous polynucleotide sequence may be capable of regulating the expression of at least one angiogenic factor of the plurality of angiogenic factors. Additionally, the at least one exogenous polynucleotide sequence may encode at least one recombinant angiogenic factor.

20        Of the plurality of fragments 66, forming micro-organs 66, at least one edge fragment is discarded and another may be automatically transferred to a viability test tube, for viability testing. Viability testing can be done for example by adding MTT to the test sample. MTT is a tetrazolium salt .

Dissolved MTT is converted into an insoluble purple formazan by cleavage of the terazolium ring by active mitochondrial dehydrogenase enzymes. The amount of color obtained is proportional to the viability and activity of the cells.

5 After rinsing and treatment, the remaining micro-organs 66 may be inserted into implanting devices 60.

Referring further to the drawings, Figures 17A and 17E, together with Figure 10A, schematically illustrates the steps in inserting micro-organs 66 into implanting devices 60, in accordance with a preferred embodiment of  
10 the present invention.

As seen in Figure 10A, a plurality of implanting devices 60 is arranged in a single line, each coupled to an micro-organ guide 54. Implanting devices 60 include slim housings 60, arranged for percutaneous insertion, protected by sterile caps 72, at their distal edges 74. They are  
15 enclosed within an implanting chamber 70, by casing 22 of device 10.

As a first step, seen in Figure 17A, sterile cap 72 is removed from each implanting device 60, exposing distal edge 74 of implanting device 60.

As a second step, seen in Figure 17B, each micro-organ guide 54, on which micro-organ 66 is held, is pulled into implanting device 60, for  
20 example, by tongues 76, so that distal edge 59 of micro-organ guide 54 protrudes from implanting device 60. Micro-organ guide 54 is pulled until position marker 68 is seen at distal edge 74 of implanting device 60.

As a second step, seen in Figures 17C and 17D, a clamp 78, within device 10 clamps micro-organ guide 54, while tongues 76 are used to break off the portion of micro-organ guide 54 distal to notch 64. The purpose of breaking off the distal portion, is to cause micro-organ 66 to be on the leading edge of micro-organ guide 54, within implanting device 60, so that leading edge will be free from guide and attach to donor tissue once partly released from implanting device 60. After the distal portion is broken off, clamp 78 releases its hold of micro-organ guide 54.

As seen in Figure 17E, implant device 60, containing micro-organ 66 and micro-organ guide 54, is detached from implanting chamber 70 of device 10, by rotation, as shown by arrow 79.

Referring further to the drawings, Figures 18A and 18C schematically illustrates the steps in implanting micro-organs 66 in a body, in accordance with a preferred embodiment of the present invention.

As a first step, shown in Figure 18A, implanting device 60 is inserted for example but not only between a muscle 84 and a skin 82 of a body, and micro-organ guide 54 is rotated by  $90^{\circ}$ , within implanting device 60. As a result, micro-organ 66 rests on muscle tissue 84.

As a second step, shown in Figure 18B, micro-organ guide 54 is pushed into implanting device 60, until a position marker 80 is no longer visible, indicating that micro-organ 66 is in its implanted position.

As a third step, shown in Figure 18C, micro-organ guide 54 is carefully pulled out, and then implanting device 60 is withdrawn.



The plurality of implanting devices 60 may be used to implant a plurality of micro-organs 66, to a subject, generally in a same area, to create a predetermined area concentration or a predetermined volume concentration of implanted micro-organs 66, in order to achieve a desired effect.

5 It will be appreciated that device 10 enables preparation of micro-organs device for immediate administration, or for storage for later use, as a sterile, functional micro-organ. The description of device 10 is given here as an example. Alternative embodiments can be envisioned that fulfill the essential features of micro-organ preparation and delivery devices.

10 It is appreciated that certain features of the invention, which are, for clarity, described in the context of separate embodiments, may also be provided in combination in a single embodiment. Conversely, various features of the invention, which are, for brevity, described in the context of a single embodiment, may also be provided separately or in any suitable  
15 subcombination.

Although the invention has been described in conjunction with specific embodiments thereof, it is evident that many alternatives, modifications and variations will be apparent to those skilled in the art. Accordingly, it is intended to embrace all such alternatives, modifications and variations that  
20 fall within the spirit and broad scope of the appended claims.

All publications, patents and patent applications mentioned in this specification are herein incorporated in their entirety by reference into the specification, to the same extent as if each individual publication, patent or

patent application was specifically and individually indicated to be incorporated herein by reference. In addition, citation or identification of any reference in this application shall not be construed as an admission that such reference is available as prior art to the present invention.

## WHAT IS CLAIMED IS:

1. A method of inducing angiogenesis in a tissue of a first mammal, the method comprising the step of implanting at least one micro-organ within the tissue of the first mammal, said at least one micro-organ being for producing a plurality of angiogenic factors and thereby inducing angiogenesis.

2. The method of claim 1, wherein said at least one micro-organ is derived from organ tissue of a second mammal.

3. The method of claim 2, wherein the first mammal and said second mammal are a single individual mammal.

4. The method of claim 2, wherein said organ is selected from the group consisting of a lung, a liver, a kidney, a muscle, a spleen a skin and a heart.

5. The method of claim 1, wherein said at least one micro-organ includes two or more cell types.

6. The method of claim 1, wherein the first mammal is a human being.

7. The method of claim 1, wherein said at least one micro-organ is cultured outside the body for at least four hours prior to implantation within the tissue of the first mammal.

8. The method of claim 1, wherein said at least one micro-organ is prepared so as to retain viability when implanted within the tissue of the first mammal.

9. The method of claim 8, wherein said at least one micro-organ has dimensions, such that cells positioned deepest within said at least one micro-organ are at least about 80 - 100 microns and not more than about 225-375 microns away from a nearest surface of said at least one micro-organ.

10. The method of claim 1, wherein each of said plurality of angiogenic factors posses a unique expression pattern within said at least one micro-organ.

11. The method of claim 1, wherein at least a portion of cells of said at least one micro-organ include at least one exogenous polynucleotide sequence selected for regulating angiogenesis.

12. The method of claim 11, wherein said at least one exogenous polynucleotide sequence is integrated into a genome of said at least a portion of said cells of said at least one micro-organ.

13. The method of claim 12, wherein said at least one exogenous polynucleotide sequence is designed for regulating expression of at least one angiogenic factor of said plurality of angiogenic factors.

14. The method of claim 13, wherein said at least one exogenous polynucleotide sequence includes an enhancer or a suppresser sequence.

15. The method of claim 11, wherein an expression product of said at least one exogenous polynucleotide sequence is capable of regulating the expression of at least one angiogenic factor of said plurality of angiogenic factors.

16. The method of claim 11, wherein said at least one exogenous polynucleotide sequence encodes at least one recombinant angiogenic factor.

17. A method of inducing angiogenesis in a tissue of a first mammal, the method comprising steps of:

(a) extracting soluble molecules from at least one micro-organ;  
and

(b) administering at least one predetermined dose of said soluble molecules extracted in step (a) into the tissue of the first mammal.

18. The method of claim 17, wherein said soluble molecules are mixed with a pharmaceutically acceptable carrier prior to step (b).

19. The method of claim 17, wherein said at least one micro-organ is derived from organ tissue of a second mammal.

20. The method of claim 17, wherein said at least one micro-organ is cultured at least four hours prior to extraction of said soluble molecules.

21. The method of claim 17, wherein said at least one micro-organ has dimensions, such that cells positioned deepest within said at least one micro-organ are at least about 80 - 100 microns and not more than about 225-375 microns away from a nearest surface of said at least one micro-organ.

22. A pharmaceutical composition comprising, as an active ingredient, a soluble molecule extract from at least one micro-organ and a pharmaceutically acceptable carrier.

23. A micro-organ comprising a plurality of cells, wherein at least a portion of said plurality of said cells including at least one exogenous

polynucleotide sequence, said at least one exogenous polynucleotide sequence being capable of regulating expression of at least one angiogenic factor expressed in said cells.

24. The micro-organ of claim 23, wherein the micro-organ is derived from organ tissue of a second mammal.

25. The micro-organ of claim 24, wherein the first mammal and said second mammal are a single individual mammal.

26. The micro-organ of claim 23, wherein said organ is selected from the group consisting of a lung, a liver, other gut derived organs, a kidney, a spleen and a heart.

27. The micro-organ of claim 23, wherein said at least one micro-organ includes two or more cell types.

28. The micro-organ of claim 23, wherein the micro-organ has dimensions, such that cells positioned deepest within the micro-organ are at least about 80 - 100 microns and not more than about 225 - 375 microns away from a nearest surface of the micro-organ.

29. The micro-organ of claim 23, wherein said at least one exogenous polynucleotide sequence is integrated into a genome of said at least a portion of said plurality of said cells.

30. The micro-organ of claim 23, wherein said at least one exogenous polynucleotide sequence includes an enhancer or a suppressor sequence.

31. The micro-organ of claim 23, wherein an expression product of said at least one exogenous polynucleotide sequence is capable of regulating the expression of said at least one angiogenic factor.

32. A method of inducing angiogenesis in a tissue of a first mammal, the method comprising the steps of:

(a) culturing at least one micro-organ in a growth medium to thereby generate a conditioned medium;

(b) collecting said conditioned medium following at least one predetermined time period of culturing; and

(c) administering at least one predetermined dose of said conditioned medium collected in step (b) into the tissue of the first mammal to thereby induce angiogenesis in the tissue.



33. The method of claim 32, wherein said at least one micro-organ is derived from organ tissue of a second mammal.

34. The method of claim 32, wherein said at least one micro-organ is cultured at least four hours prior to collection of said conditioned medium.

35. The method of claim 32, wherein said at least one micro-organ has dimensions, such that cells positioned deepest within said at least one micro-organ are at least about 80 - 100 microns and not more than about 225-375 microns away from a nearest surface of said at least one micro-organ.

36. The method of claim 32, wherein said growth medium is a minimal essential medium.

37. An apparatus for generating micro-organs from a tissue biopsy and for administering the micro-organs into a subject, the apparatus comprising:

(a) a cutting chamber for cutting the tissue biopsy into a plurality of micro-organs; and

(b) an implanting mechanism for administering the plurality of micro-organs into the subject, said implanting mechanism being operably coupled to said cutting chamber.

38. The apparatus of claim 37, wherein said cutting chamber has an inlet/outlet for introducing and removing reagents.

39. The apparatus of claim 37, wherein said cutting chamber has an inlet for introducing the tissue biopsy therein.

40. The apparatus of claim 37, further comprising a viability testing chamber operably coupled to said cutting chamber for testing a viability of at least one sacrificial micro-organ of said plurality of micro-organs.

41. The apparatus of claim 37, wherein said implanting mechanism comprises a multi-channel planter and corresponding advancing elements for advancing said plurality of micro-organs from said cutting chamber to said multi-channel planter and further for administering the plurality of micro-organs into the subject.

42. The apparatus of claim 37, further comprising a processing chamber being operably coupled to said cutting chamber and said implanting mechanism for processing said micro-organs prior to said administering.

43. The apparatus of claim 42, wherein said processing chamber has an inlet/outlet for introducing and removing processing reagents.

44. The apparatus of claim 37, wherein said cutting chamber is designed and constructed such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ.

45. The apparatus of claim 37, wherein said cutting chamber comprises a cutting mechanism having a plurality of blades movable to cut the tissue biopsy into said plurality of micro-organs.

46. The apparatus of claim 45, wherein said blades are so disposed with respect to one another such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80-100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ.

47. The apparatus of claim 45, wherein each of said plurality of blades has a translatable angled cutting edge.

48. The apparatus of claim 45, wherein each of said plurality of blades is a rotatable disc-blade.

49. An apparatus for generating micro-organs from a tissue biopsy, the apparatus comprising:

(a) a cutting chamber for cutting the tissue biopsy into a plurality of micro-organs; and

(b) a viability testing chamber operably coupled to said cutting chamber for testing a viability of at least one sacrificial micro-organ of said plurality of micro-organs.

50. The apparatus of claim 49, wherein said cutting chamber has an inlet/outlet for introducing and removing reagents:

51. The apparatus of claim 49, wherein said cutting chamber has an inlet for introducing the tissue biopsy therein.

52. The apparatus of claim 49, wherein said cutting chamber is designed and constructed such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ.

53. The apparatus of claim 49, wherein said cutting chamber comprises a cutting mechanism having a plurality of blades movable to cut the tissue biopsy into said plurality of micro-organs.

54. The apparatus of claim 53, wherein said blades are so disposed with respect to one another such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225 - 375 microns away from a nearest surface of said micro-organ.

55. The apparatus of claim 53, wherein each of said plurality of blades has a translatable angled cutting edge.

56. The apparatus of claim 53, wherein each of said plurality of blades is a rotatable disc-blade.

57. An apparatus for generating micro-organs from a tissue biopsy,  
the apparatus comprising:

(a) a cutting chamber for cutting the tissue biopsy into a plurality of micro-organs;

(b) a processing chamber being operably coupled to said cutting chamber for processing said micro-organs; and

(c) an advancing mechanism for advancing said micro-organs from said cutting chamber into said processing chamber.

58. The apparatus of claim 57, wherein said processing chamber has an inlet/outlet for introducing and removing processing reagents.

59. The apparatus of claim 57, wherein said cutting chamber has an inlet/outlet for introducing and removing reagents.

60. The apparatus of claim 57, wherein said cutting chamber has an inlet for introducing the tissue biopsy therein.

61. The apparatus of claim 57, wherein said cutting chamber is designed and constructed such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ.

62. The apparatus of claim 57, wherein said cutting chamber comprises a cutting mechanism having a plurality of blades movable to cut the tissue biopsy into said plurality of micro-organs.

63. The apparatus of claim 62, wherein said blades are so disposed with respect to one another such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ.

64. The apparatus of claim 62, wherein each of said plurality of blades has a translatable angled cutting edge.

65. The apparatus of claim 62, wherein each of said plurality of blades is a rotatable disc-blade.

66. A method of generating micro-organs from a tissue biopsy and for administering the micro-organs into a subject, the method comprising:

providing an apparatus which comprises:

(a) a cutting chamber for cutting the tissue biopsy into a plurality of micro-organs; and

(b) an implanting mechanism for administering the plurality of micro-organs into the subject, said implanting mechanism being operably coupled to said cutting chamber.

placing the tissue biopsy in said cutting chamber and cutting the tissue biopsy into the plurality of micro-organs; and

using said implanting mechanism for administering the plurality of micro-organs into the subject.

67. The method of claim 66, wherein the micro-organs serve as angiopumps.

68. The method of claim 66, wherein said cutting chamber has an inlet/outlet for introducing and removing reagents, the method further comprising washing said micro-organs in said cutting chamber prior to using said implanting mechanism for administering the plurality of micro-organs into the subject.

69. The method of claim 66, wherein said cutting chamber has an inlet for introducing the tissue biopsy therein, the method comprising placing the tissue biopsy in said cutting chamber through said inlet.

70. The method of claim 66, wherein said apparatus further comprises a viability testing chamber operably coupled to said cutting chamber for testing a viability of at least one sacrificial micro-organ of said plurality of micro-organs, the method further comprising testing said viability of said at least one sacrificial micro-organ of said plurality of micro-organs prior to using said implanting mechanism for administering the plurality of micro-organs into the subject.



71. The method of claim 66, wherein said implanting mechanism comprises a multi-channel implanter and corresponding advancing elements for advancing said plurality of micro-organs from said cutting chamber to said multi-channel implanter and further for administering the plurality of micro-organs into the subject, the method comprising administering the plurality of micro-organs into the subject using said advancing elements.

72. The method of claim 66, further comprising a processing chamber being operably coupled to said cutting chamber and said administration mechanism for processing said micro-organs prior to said administering, the method further comprising processing said micro-organs prior to said administering.

73. The method of claim 72, wherein said processing said micro-organs prior to said administering comprises at least one a process selected from the group consisting of washing, transforming, culturing, and a combination thereof.

74. The method of claim 72, wherein said processing said micro-organs prior to said administering comprises culturing for at least one hour.

75. The method of claim 72, wherein said processing said micro-organs prior to said administering comprises transforming by introducing to

at least a portion of cells of said micro-organs at least one exogenous polynucleotide sequence selected for regulating angiogenesis.

76. The method of claim 75, wherein said at least one exogenous polynucleotide sequence is integrated into a genome of said at least said portion of said cells of said micro-organs.

77. The method of claim 76, wherein said at least one exogenous polynucleotide sequence is designed for regulating expression of at least one angiogenic factor of said plurality of angiogenic factors.

78. The method of claim 77, wherein said at least one exogenous polynucleotide sequence includes an enhancer or a suppresser sequence.

79. The method of claim 75, wherein an expression product of said at least one exogenous polynucleotide sequence is capable of regulating the expression of at least one angiogenic factor of said plurality of angiogenic factors.

80. The method of claim 75, wherein said at least one exogenous polynucleotide sequence encodes at least one recombinant angiogenic factor.

81. The method of claim 72, wherein said processing chamber has an inlet/outlet for introducing and removing processing reagents, the method comprising introducing at least one processing reagent into said processing chamber through said inlet/outlet.

82. The method of claim 66, wherein said cutting chamber is designed and constructed such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ, the method further comprising using said cutting chamber to cut the tissue biopsy into said plurality of micro-organs each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from said nearest surface of said micro-organ.

83. The method of claim 66, wherein said cutting chamber comprises a cutting mechanism having a plurality of blades movable to cut the tissue biopsy into said plurality of micro-organs, the method comprising using said plurality of blades to cut the tissue biopsy into said plurality of micro-organs.

84. The method of claim 83, wherein said blades are so disposed with respect to one another such that once the tissue biopsy is cut into said plurality of micro-organs, each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from a nearest surface of said micro-organ, the method comprising using said plurality of blades to cut the tissue biopsy into said plurality of micro-organs each of said micro-organs such that cells positioned deepest within a micro-organ of said plurality of micro-organs are at least about 80 - 100 microns and not more than 225-375 microns away from said nearest surface of said micro-organ.

85. The method of claim 83, wherein each of said plurality of blades has a translatable angled cutting edge, the method comprising translating said angled cutting edge with respect to the tissue biopsy, so as to cut the tissue biopsy into said plurality of micro-organs.

86. The method of claim 83, wherein each of said plurality of blades is a rotatable disc-blade, the method comprising moving said rotatable disc-blade with respect to the tissue biopsy, so as to cut the tissue biopsy into said plurality of micro-organs.

87. The method of claim 66, wherein the tissue biopsy is derived from a tissue or organ selected from the group consisting of lung, liver, kidney, muscle, spleen, skin, heart, lymph node and bone marrow.

88. The method of claim 66, wherein a donor of the tissue biopsy and the subject are the same individual.

89. The method of claim 66, wherein a donor of the tissue biopsy and the subject are different individuals.

90. The method of claim 66, wherein a donor of the tissue biopsy is a human.

91. The method of claim 66, wherein a donor of the tissue biopsy is a non-human mammal.

92. The method of claim 66, wherein the subject is a non-human mammal.

93. The method of claim 66, wherein the subject is a human.

94. The method of claim 66, wherein administering the plurality of micro-organs into the subject is effected via transmucosal or parenteral administration routes.

95. The method of claim 94, wherein said transmucosal or parenteral administration routes are selected from the group consisting of intramuscular, subcutaneous, intramedullary, intrathecal, direct intraventricular, intravenous, intraperitoneal, intranasal and intraocular administration routes.

96. A device for micro-organ preparation and delivery, comprising:  
a tissue cutter, for cutting a tissue biopsy into a plurality of fragments, forming a plurality of micro-organs: and

at least one implanting device, detachably coupled to said tissue cutter, for receiving a micro-organ, of said plurality of micro-organs, when coupled to said tissue cutter, and for implanting said micro-organ into a subject, after decoupling from said tissue cutter.

97. The device of claim 96, further comprising a tissue scraper, for obtaining said tissue biopsy.

98. The device of claim 97, wherein said tissue scraper is adapted for preparing said tissue biopsy to a predetermined width.

99. The device of claim 97, wherein said tissue scraper is adapted for preparing said tissue biopsy to a predetermined length.

100. The device of claim 97, wherein said tissue scraper is adapted for preparing said tissue biopsy to a predetermined thickness.

101. The device of claim 97, wherein said tissue scraper has a replaceable blade.

102. The device of claim 96, wherein said device is sealed within a base, a ramp, and a casing.

103. The device of claim 96, wherein said device includes a control system.

104. The device of claim 96, wherein said device includes at least one automated travel mechanism for transferring the tissue biopsy from one region of said device to another.

105. The device of claim 96, wherein said device includes a washing apparatus for rinsing the tissue biopsy.

106. The device of claim 105, wherein said washing apparatus is further operative for applying a medium to the tissue biopsy.

107. The device of claim 96, wherein said device is further operative as a tissue treatment chamber.

108. The device of claim 96, wherein said device includes apparatus for controlling the temperature therein.

109. The device of claim 96, wherein said tissue cutter comprises a plurality of parallel, surgical-grade blades, designed to cut the tissue biopsy into said plurality of fragments, forming said micro-organs, such that cells positioned deepest within any one of said micro-organs are at least about 80 – 100 microns and not more than about 225 - 375 microns away from a nearest surface.

110. The device of claim 96, wherein said tissue cutter comprises a plurality of parallel surgical-grade blades, arranged at an angle to the tissue biopsy.

111. The device of claim 96, wherein said tissue cutter comprises a plurality of parallel surgical-grade blades, arranged as rotatable disc-blades.



112. The device of claim 96, wherein said device comprises a viability testing chamber for testing a viability of at least one micro-organ of said plurality of micro-organs.

113. The device of claim 96, wherein said tissue cutter is operative to cut the tissue biopsy, to form said micro-organs, and to arrange each of said micro-organs on a single guide of a plurality of guides, in a single operation.

114. The device of claim 113, wherein said at least one implanting device includes a slim housing, adapted for percutaneous insertion, and operable to receive one of said plurality of guides.

115. The device of claim 113, wherein said at least one implanting device includes a plurality of implanting devices, each operable to receive one of said plurality of guides.

116. The device of claim 113, wherein each of said plurality of micro-organ guides includes a position marker for indicating when said micro-organ, arranged on it, is positioned for implanting.

117. The device of claim 113, wherein each of said micro-organ guides includes a notch for breaking off a distal portion thereof, to allow said micro-organ, arranged on it, to form a leading edge.

118. The device of claim 113, wherein each of said plurality of micro-organ guides includes a position marker for indicating when said micro-organ, arranged on it, is implanted.

119. The device of claim 96, wherein said device is disposable.

120. A method for micro-organ preparation and delivery, comprising:

scraping a tissue biopsy;

cutting the tissue biopsy to a plurality of fragments, forming a plurality of micro-organs; and

implanting at least one of said plurality of micro-organs.

121. The method of claim 120, wherein said micro-organ serves as an angiopump.

122. The method of claim 120, and further including treating the tissue biopsy, prior to implanting.

123. The method of claim 122, wherein said treating is selected from the group consisting of washing, transforming, culturing, and a combination thereof.

124. The method of claim 120, wherein:

said cutting further includes cutting to a first plurality of tissue fragments, forming a first plurality of micro-organs; and

said implanting further includes implanting a second plurality of fragments, wherein said second plurality is smaller than said first plurality by at least one,

wherein said method further includes using at least one of said first plurality of tissue fragments for a viability test.

125. The method of claim 120, wherein said cutting includes cutting the tissue biopsy into said plurality of fragments, forming said micro-organs, such that cells positioned deepest within any one of said micro-organs are at least about 80 microns and not more than about 375 microns away from a nearest surface.

126. The method of claim 120, wherein said implanting further includes implanting a plurality of micro-organs within a preselected area of said subject, for a predetermined area concentration of micro-organs.

127. The method of claim 120, wherein said implanting further includes implanting a plurality of micro-organs within a preselected volume of said subject, for a predetermined volume concentration of micro-organs.

128. A method for micro-organ preparation and delivery, comprising:

employing a device for micro-organ preparation and delivery, which includes:

a tissue scraper, for obtaining a tissue biopsy;

a tissue cutter, for cutting the tissue biopsy into a plurality of fragments, forming a plurality of micro-organs;  
and

at least one implanting device, detachably coupled to said tissue cutter, for receiving a micro-organ, of said plurality of micro-organs, when coupled to said tissue cutter, and for implanting said micro-organ into a subject, after decoupling from said tissue cutter;

scraping the tissue biopsy, with said tissue scraper;

cutting the tissue biopsy to said plurality of fragments, forming said plurality of micro-organs, with said tissue cutter;

mounting said micro-organ, of said plurality of micro-organs, on said at least one implanting device;

decoupling said at least one implanting device; and

implanting said micro-organ, with said at least one implanting device.

129. The method of claim 128, wherein said micro-organ serves as an angiopump.

130. The method of claim 128, wherein said device is sealed within a base, a ramp, and a casing.

131. The method of claim 128, wherein said device includes at least one automated travel mechanism for transferring the tissue biopsy from one region of said device to another.

132. The method of claim 128, wherein said tissue scraper is adapted for scraping said tissue to a predetermined width.

133. The method of claim 128, wherein said tissue scraper is adapted for scraping said tissue to a predetermined length.

134. The method of claim 128, wherein said tissue scraper is adapted for scraping said tissue to a predetermined thickness.

135. The method of claim 128, wherein said tissue scraper has a replaceable blade.

136. The method of claim 128, wherein said device includes a washing apparatus for rinsing the tissue biopsy.

137. The method of claim 128, wherein said washing apparatus is further operative for applying a medium onto the tissue biopsy.

138. The method of claim 128, and further including treating the tissue biopsy, prior to implanting.

139. The method of claim 138, wherein said treating is selected from the group consisting of washing, transforming, culturing, and a combination thereof.

140. The method of claim 128, wherein said device includes apparatus for controlling the temperature therein.

141. The method of claim 128, wherein said tissue cutter comprises a plurality of parallel, surgical-grade blades, designed to cut the tissue biopsy into said plurality of fragments, forming said micro-organs, such that cells positioned deepest within any one of said micro-organs are at least about 80 – 100 microns and not more than about 225 – 375 microns away from a nearest surface.

142. The method of claim 128, wherein said tissue cutter comprises a plurality of parallel surgical-grade blades, arranged at an angle to the tissue biopsy.

143. The method of claim 128, wherein said tissue cutter comprises a plurality of parallel surgical-grade blades, arranged as rotatable disc-blades.

144. The method of claim 128, wherein said device comprises a viability testing chamber for testing a viability of at least one micro-organ of said plurality of micro-organs.

145. The method of claim 128, wherein said cutting further includes arranging each of said micro-organs on a single guide of a plurality of guides.

146. The method of claim 145, wherein said at least one implanting device includes a slim housing, adapted for percutaneous insertion, and operable to receive one of said plurality of guides.

147. The method of claim 145, wherein said at least one implanting device includes a plurality of implanting devices, each operable to receive one of said plurality of guides.

148. The method of claim 145, wherein each of said plurality of micro-organ guides includes a position marker for indicating when said micro-organ, arranged on it, is positioned for implanting.

149. The method of claim 145, wherein each of said micro-organ guides includes a notch for breaking off a distal portion thereof, to allow said micro-organ, arranged on it, to form a leading edge.

150. The method of claim 145, wherein each of said plurality of micro-organ guides includes a position marker for indicating when said micro-organ, arranged on it, is implanted.

151. The method of claim 128, wherein said method further includes disposing said device after one use.

152. The method of claim 128, wherein the tissue biopsy is derived from a tissue or organ selected from the group consisting of lung, liver, kidney, muscle, spleen, skin, heart, lymph node and bone marrow.

153. The method of claim 128, wherein a donor of the tissue biopsy and the subject are the same individual.



154. The method of claim 128, wherein a donor of the tissue biopsy and the subject are different individuals.
155. The method of claim 128, wherein a donor of the tissue biopsy is a human.
156. The method of claim 128, wherein a donor of the tissue biopsy is a non-human mammal.
157. The method of claim 128, wherein the subject is a non-human mammal.
158. The method of claim 128, wherein the subject is a human.
159. The method of claim 128, wherein said device includes a control system.
160. The method of claim 128, wherein:  
said cutting further includes cutting to a first plurality of tissue fragments, forming a first plurality of micro-organs; and  
said implanting further includes implanting a second plurality of micro-organs, wherein said second plurality is selected from the group consisting of a plurality which is equal to said first plurality, a plurality

which is smaller than said second plurality by one, and a plurality which is smaller than said second plurality by two.

161. The method of claim 128, wherein:

said cutting further includes cutting to a first plurality of tissue fragments, forming a first plurality of micro-organs; and

said implanting further includes implanting a second plurality of fragments,

wherein said second plurality is smaller than said first plurality by one,

and wherein said method further includes using an edge fragment for a viability test.

162. The method of claim 128, wherein:

said cutting further includes cutting to a first plurality of tissue fragments, forming a first plurality of micro-organs; and

said implanting further includes implanting a second plurality of tissue fragments, wherein said second plurality is smaller than said first plurality by two,

wherein said method further includes:

using a first edge fragment for a viability test; and

discarding a second edge fragment.

163. The method of claim 128, wherein said cutting includes cutting the tissue biopsy into said plurality of fragments, forming said micro-organs, such that cells positioned deepest within any one of said micro-organs are at least about 80 microns and not more than about 375 microns away from a nearest surface.

164. The method of claim 128, wherein said cutting includes cutting the tissue biopsy into said plurality of fragments, forming said micro-organs, such that cells positioned deepest within any one of said micro-organs are at least about 100 microns and not more than about 225 microns away from a nearest surface.

165. The method of claim 128, wherein said implanting further includes implanting a plurality of micro-organs within a preselected area of said subject, for a predetermined area concentration of micro-organs.

166. The method of claim 128, wherein said implanting further includes implanting a plurality of micro-organs within a preselected volume of said subject, for a predetermined volume concentration of micro-organs.

167. The method of claim 128, wherein said tissue biopsy is a split-thickness tissue biopsy.

# ABSTRACT OF THE DISCLOSURE

A method, extract, and pharmaceutical composition for inducing angiogenesis in a tissue of a mammal, and of a device for the preparation and delivery of micro-organs into a mammal, are provided.

Docket No.  
02/23451**Declaration and Power of Attorney For Patent Application****English Language Declaration**

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

**METHOD AND DEVICE FOR INDUCING  
BIOLOGICAL PROCESSES BY MICRO-ORGANS**

the specification of which



is attached hereto.



was filed on \_\_\_\_\_ as United States Application No. or PCT

International Application Number \_\_\_\_\_

and was amended on \_\_\_\_\_

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose to the United States Patent and Trademark Office all information known to me to be material to patentability as defined in Title 37, Code of Federal Regulations, Section 1.56.

I hereby claim foreign priority benefits under Title 35, United States Code, Section 119(a)-(d) or Section 365(b) of any foreign application(s) for patent or inventor's certificate, or Section 365(a) of any PCT International application which designated at least one country other than the United States, listed below and have also identified below, by checking the box, any foreign application for patent or inventor's certificate or PCT International application having a filing date before that of the application on which priority is claimed.

Prior Foreign Application(s)

Priority Not Claimed

\_\_\_\_\_  
(Number)

\_\_\_\_\_  
(Country)

\_\_\_\_\_  
(Day/Month/Year Filed)



\_\_\_\_\_  
(Number)

\_\_\_\_\_  
(Country)

\_\_\_\_\_  
(Day/Month/Year Filed)



\_\_\_\_\_  
(Number)

\_\_\_\_\_  
(Country)

\_\_\_\_\_  
(Day/Month/Year Filed)



I hereby claim the benefit under 35 U.S.C. Section 119(e) of any United States provisional application(s) listed below:

<u>60/140,748</u>	<u>25 June 1999</u>
(Application Serial No.)	(Filing Date)
(Application Serial No.)	(Filing Date)
(Application Serial No.)	(Filing Date)

I hereby claim the benefit under 35 U.S.C. Section 120 of any United States application(s), or Section 365(c) of any PCT International application designating the United States, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of 35 U.S.C. Section 112. I acknowledge the duty to disclose to the United States Patent and Trademark Office all the information known to me to be material to patentability as defined in Title 37, C.F.R., Section 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application:

<u>10/009,520</u>	<u>22 June 2000</u>	<u>Pending</u>
(Application Serial No.)	(Filing Date)	(Status)
		(patented, pending, abandoned)
<u>PCT/IL00/00365</u>	<u>22 June 2000</u>	 
(Application Serial No.)	(Filing Date)	(Status)
		(patented, pending, abandoned)
(Application Serial No.)	(Filing Date)	(Status)
		(patented, pending, abandoned)

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. *(list name and registration number)*

**SOL SHEINBEIN, Registration Number 25,457**  
**MARTIN MOYNIHAN, Registration Number 40,338**  
**ROCHEL ABBOUDI, Registration Number 44,490**

Send Correspondence to:

**G.E. EHRLICH (1995) LTD.**  
**c/o ANTHONY CASTORINA**  
**2001 JEFFERSON DAVIS HIGHWAY**  
**SUITE 207**  
**ARLINGTON, VIRGINIA 22202, USA**

Direct Telephone Calls to: *(name and telephone number)*

**Anthony Castorina**

**Tel. No. (703) 415-1581**  
**Fax No. (703) 415- 4864**


FULL NAME OF SOLE OR FIRST INVENTOR	<b>Eduardo N. MITRANI</b>	
Sole or first inventor's signature		Date <u>July 9, 2002</u>
Residence	: 22 Reuven Street, 93510 Jerusalem, Israel	
Citizenship	: Israeli	
Post Office Address	: 22 Reuven Street, 93510 Jerusalem, Israel	



FIG.1

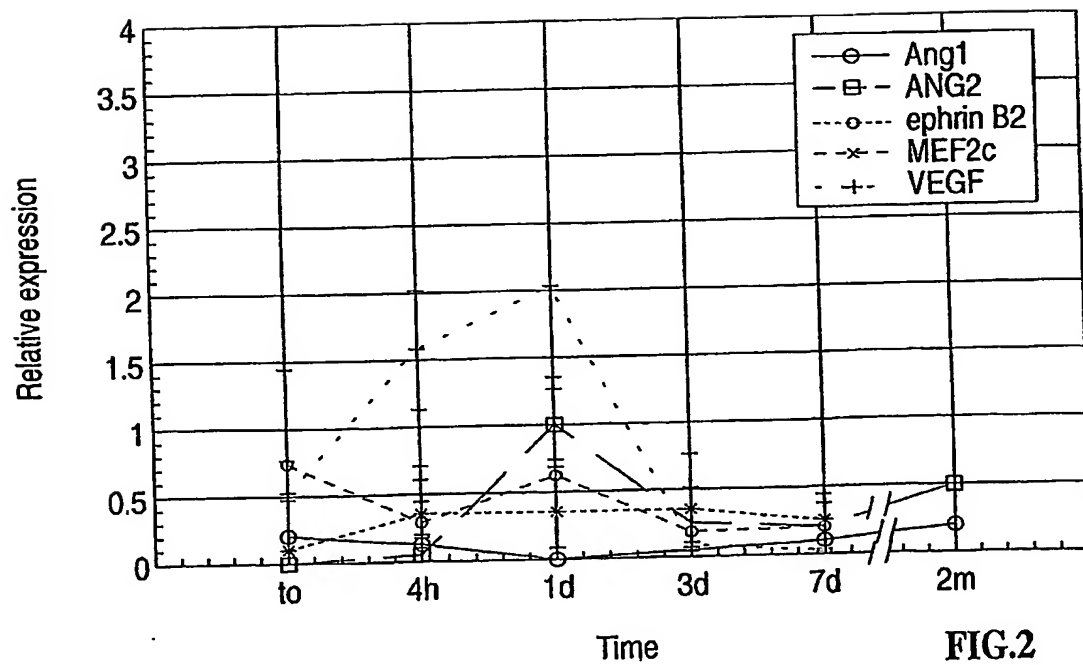


FIG.2

BEST AVAILABLE COPY



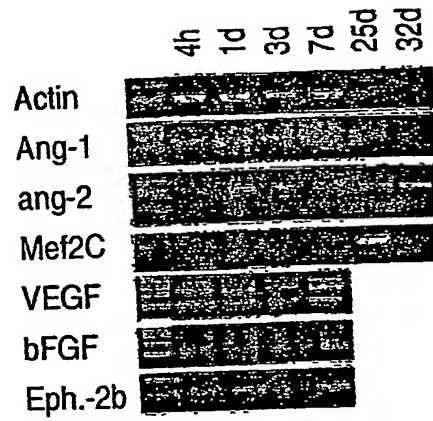


FIG.3

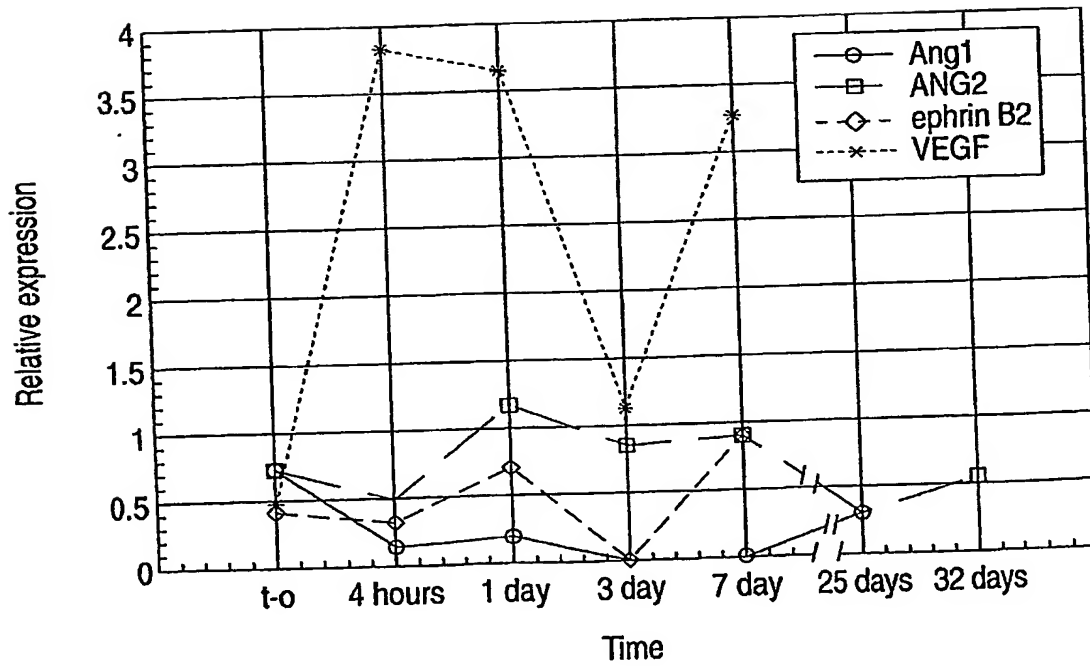


FIG.4

BEST AVAILABLE COPY

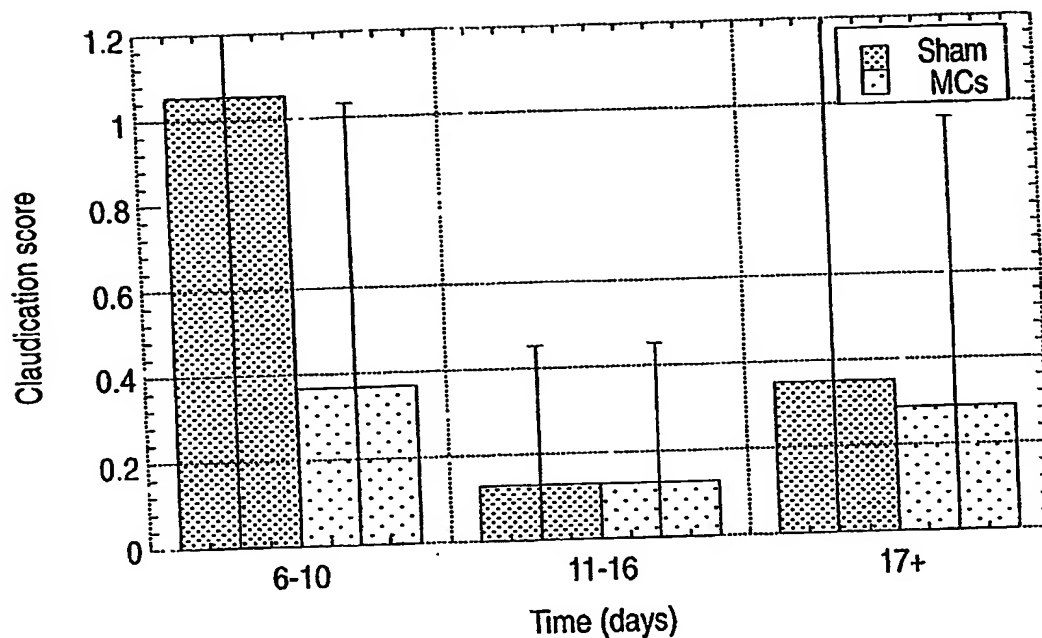


FIG.5

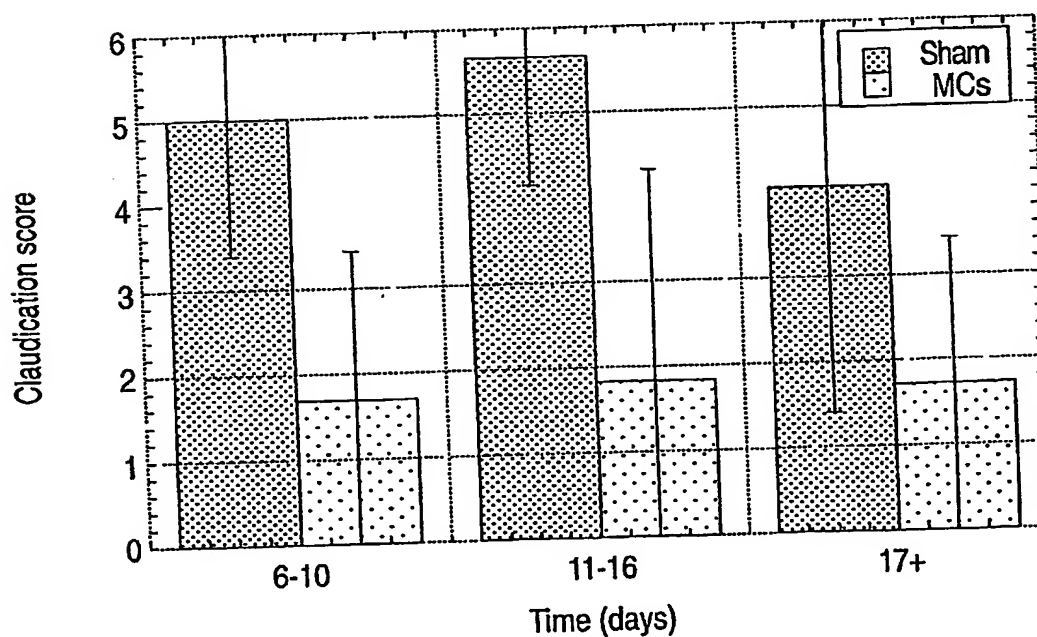
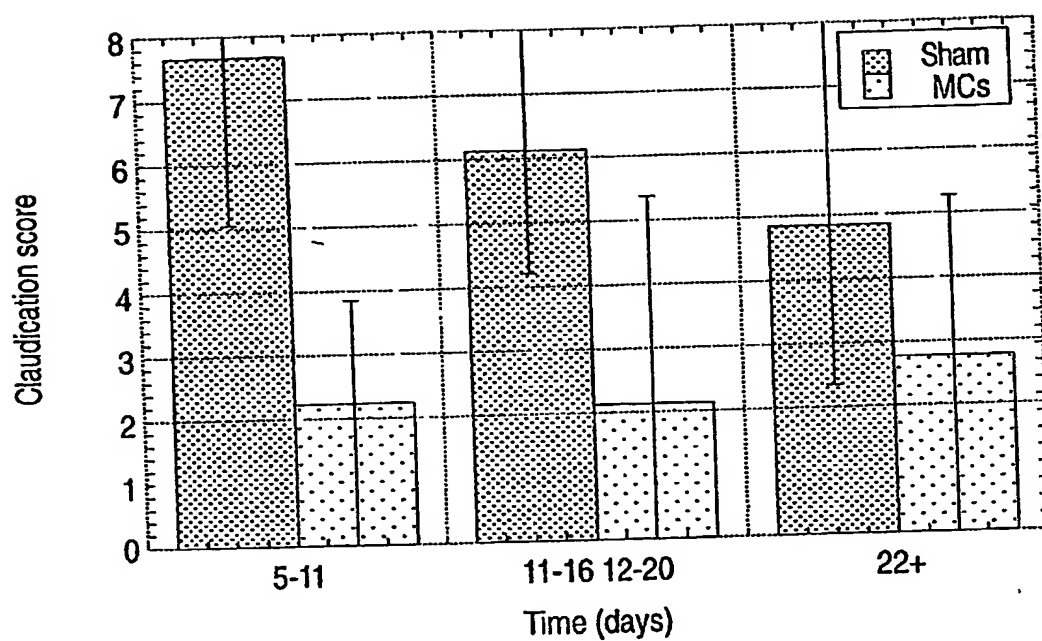


FIG.6

BEST AVAILABLE COPY

**FIG.7****BEST AVAILABLE COPY**

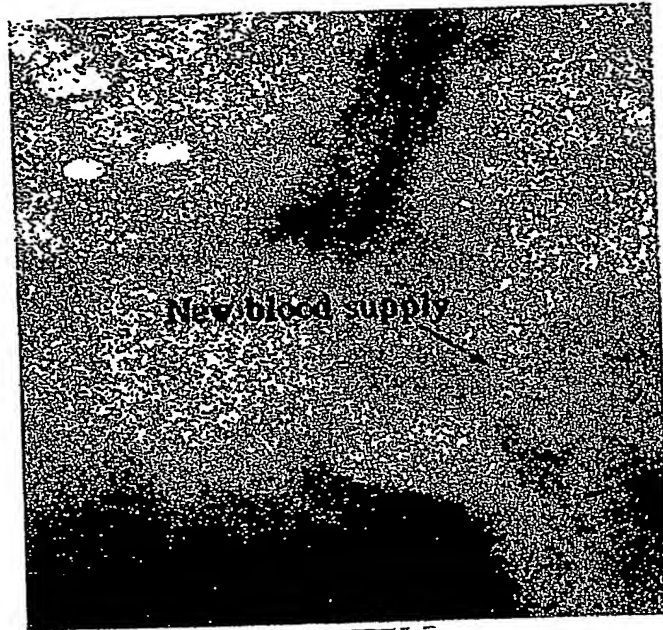


FIG.8

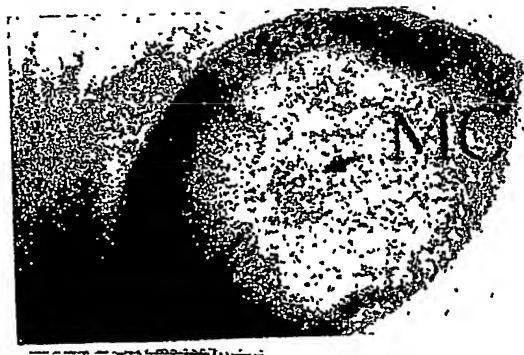


FIG.9

REST AVAILABLE COPY

Figure 10 A

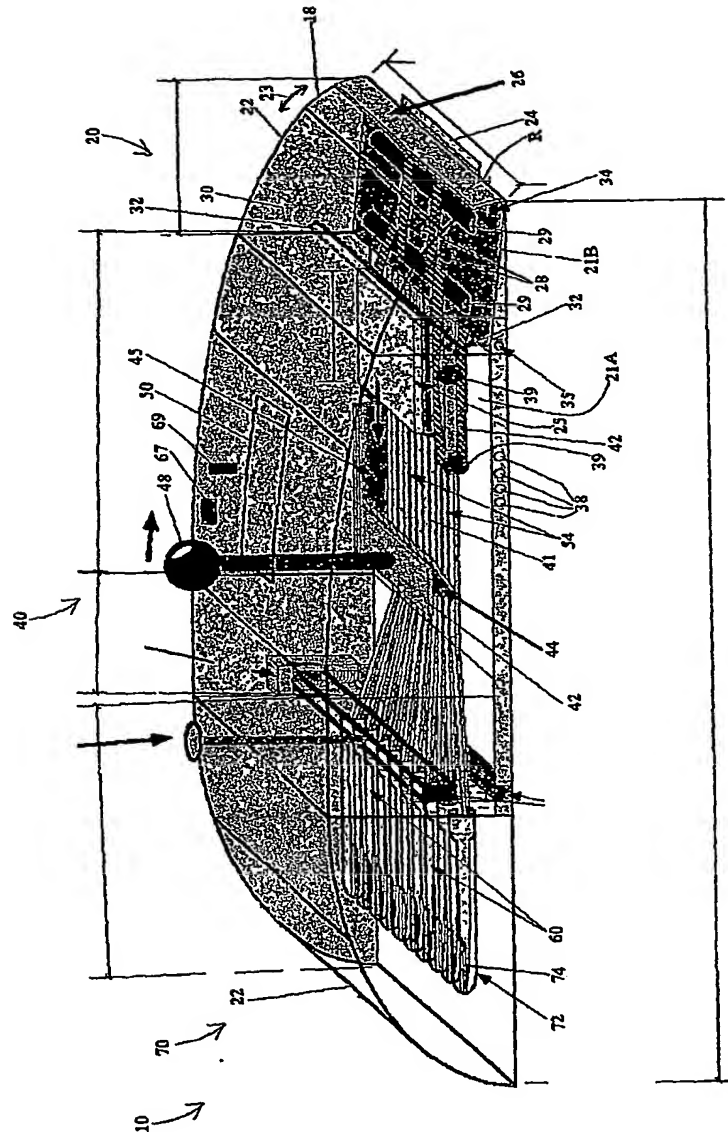
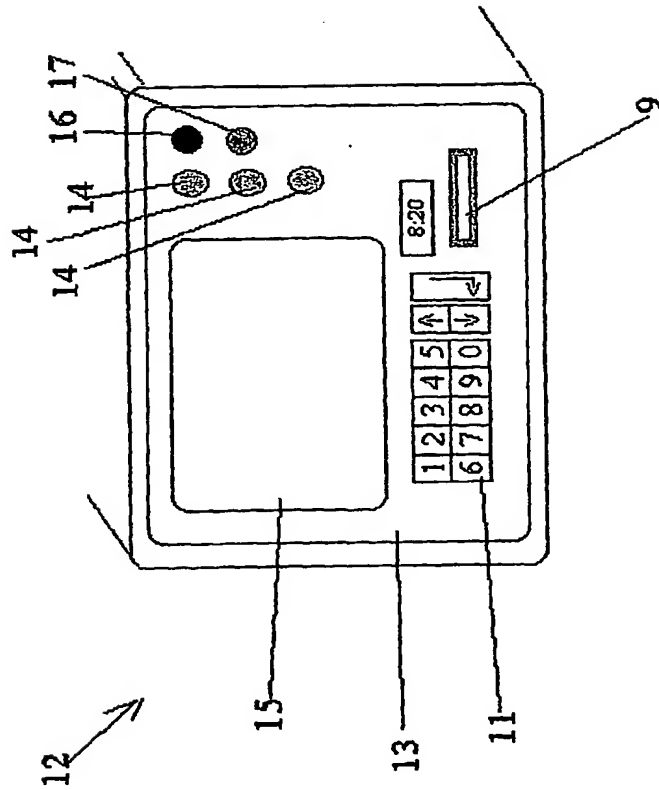


Figure 10B



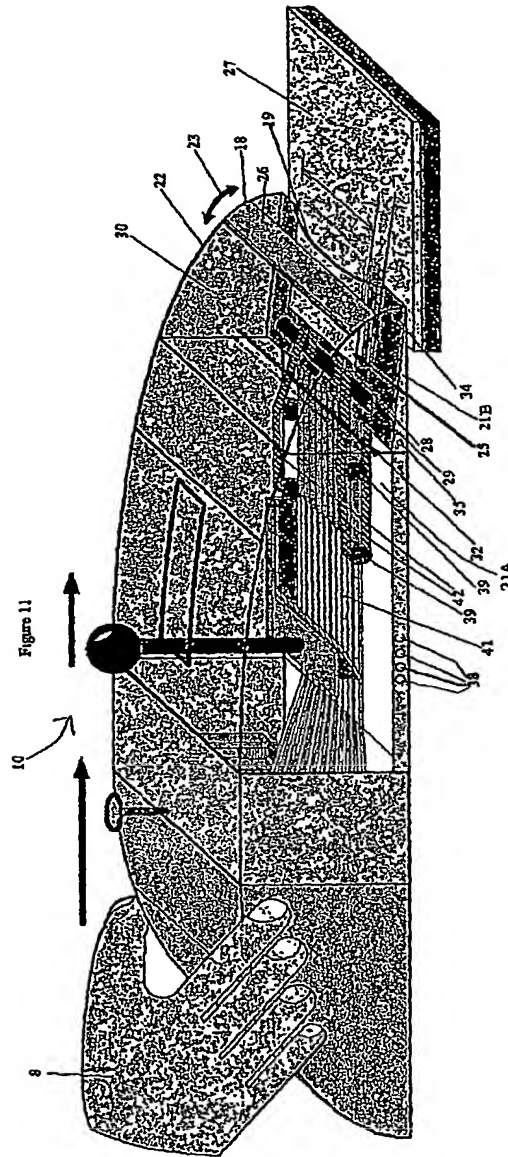
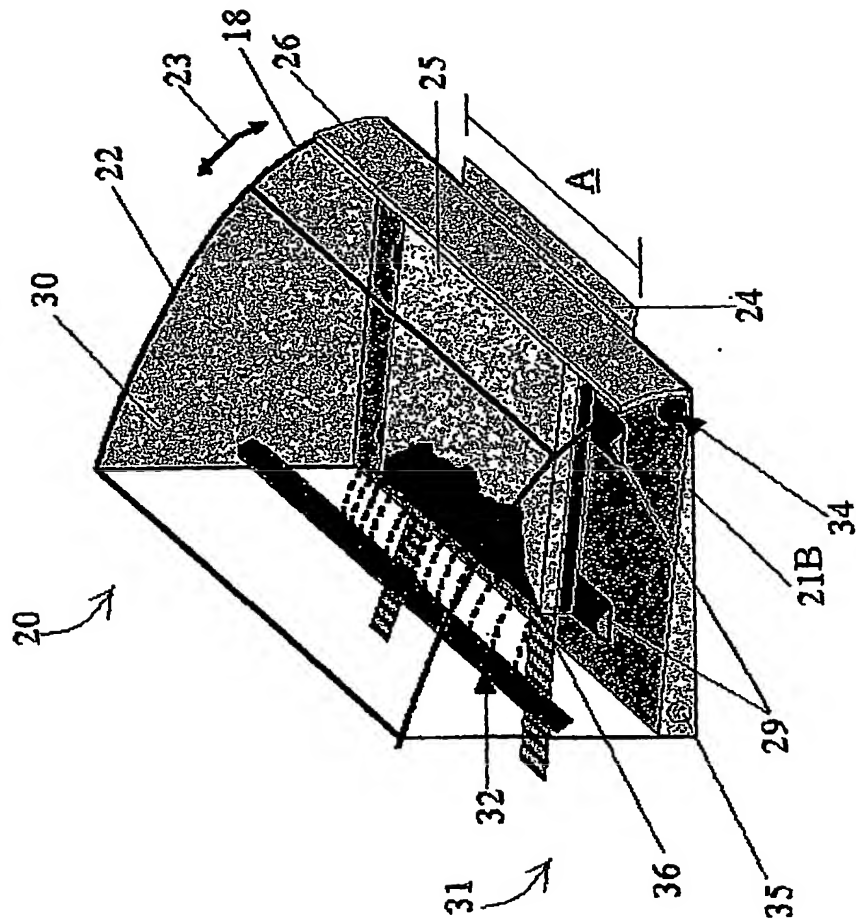


Figure 12





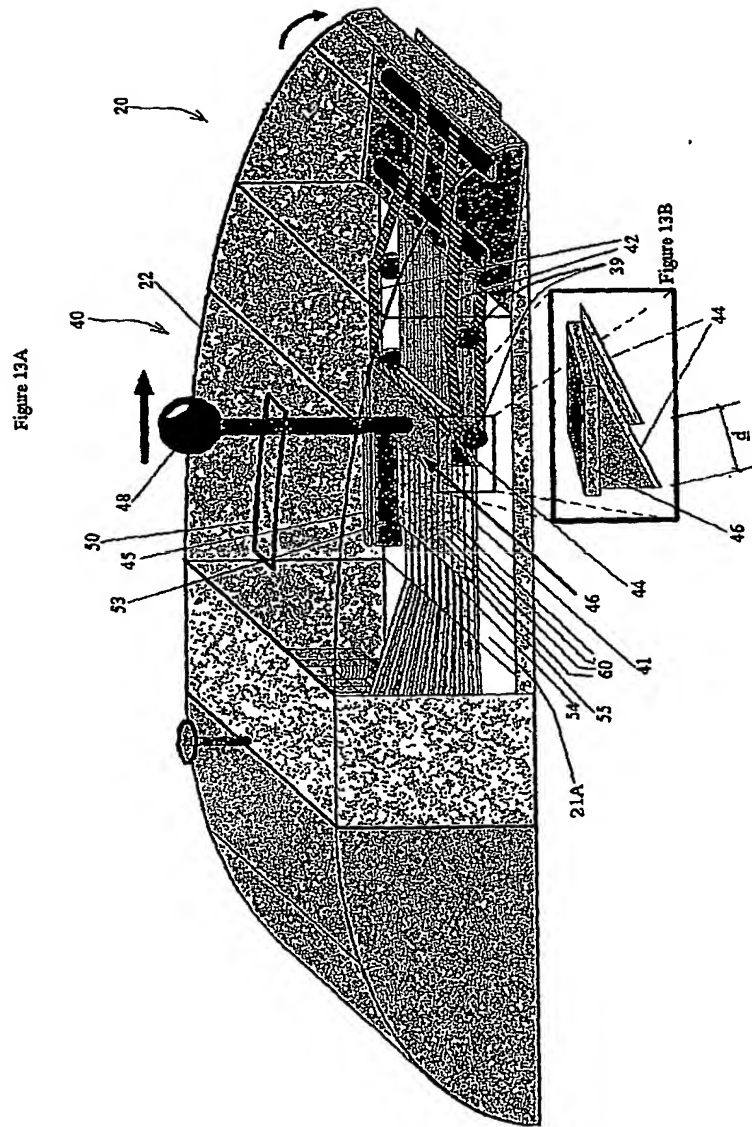


Figure 14A

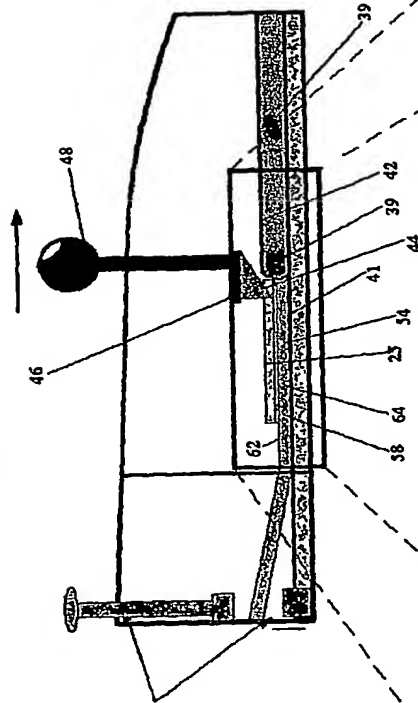


Figure 14B

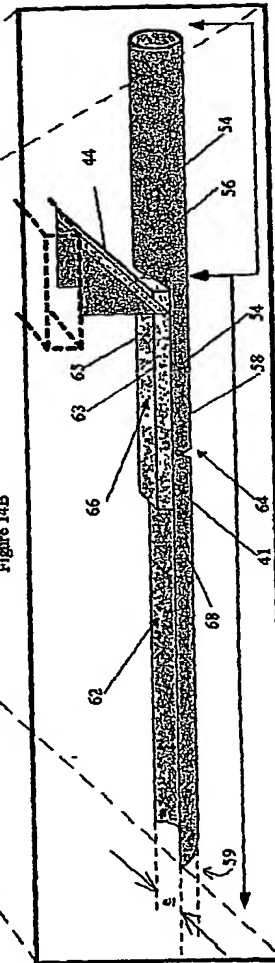


Figure 14D is a perspective view of a rectangular panel 25. The panel has a central area 53 and a surrounding border 52. The border 52 is composed of a top layer 57 and a bottom layer 55. Arrows indicate the direction of light or signal passing through the panel.

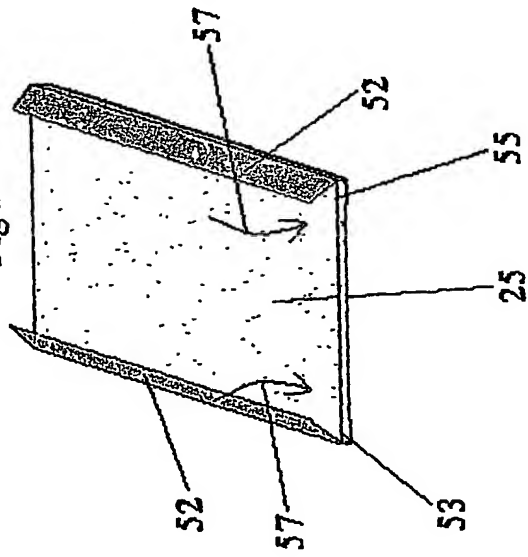
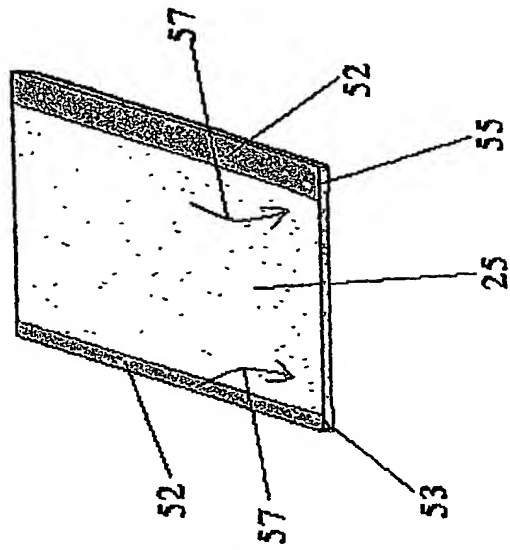


Figure 15

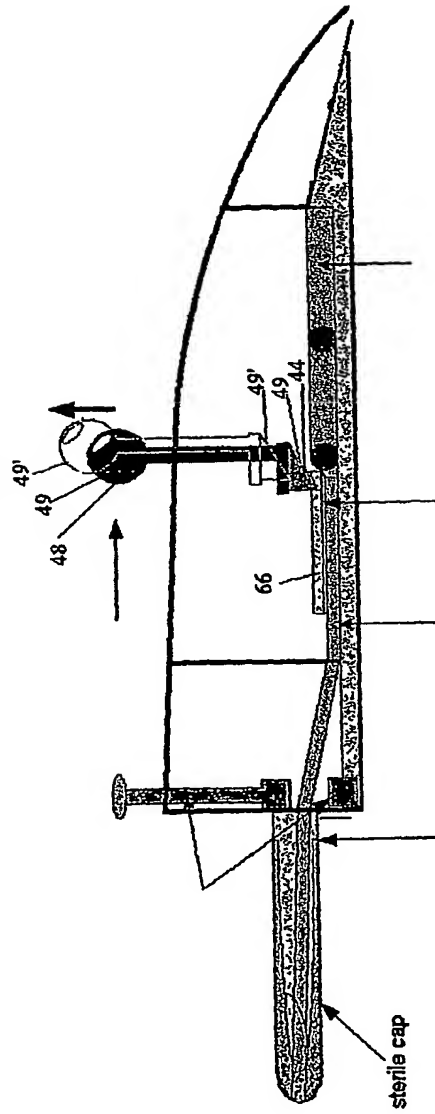


Figure 16A

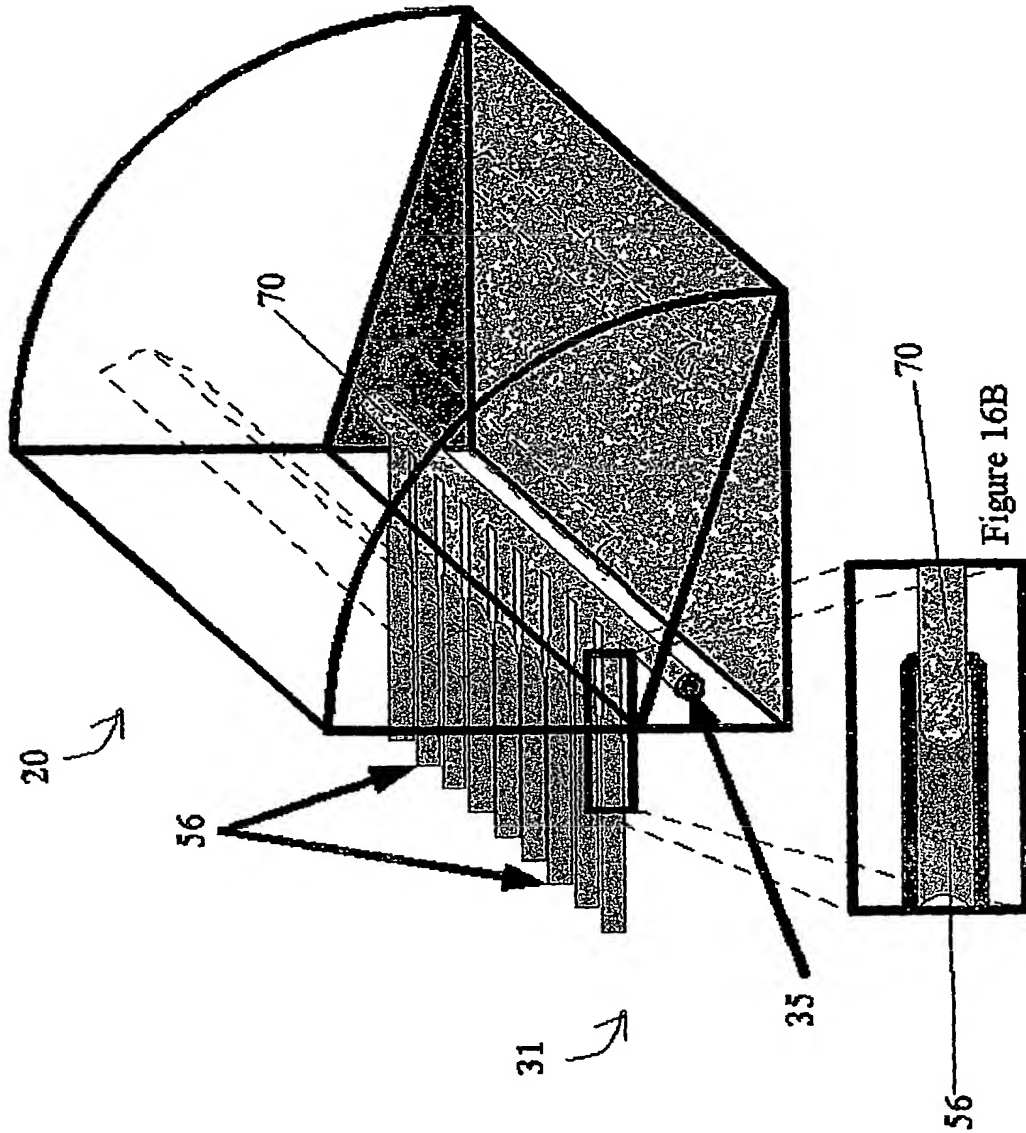


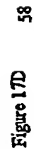
Figure 16B

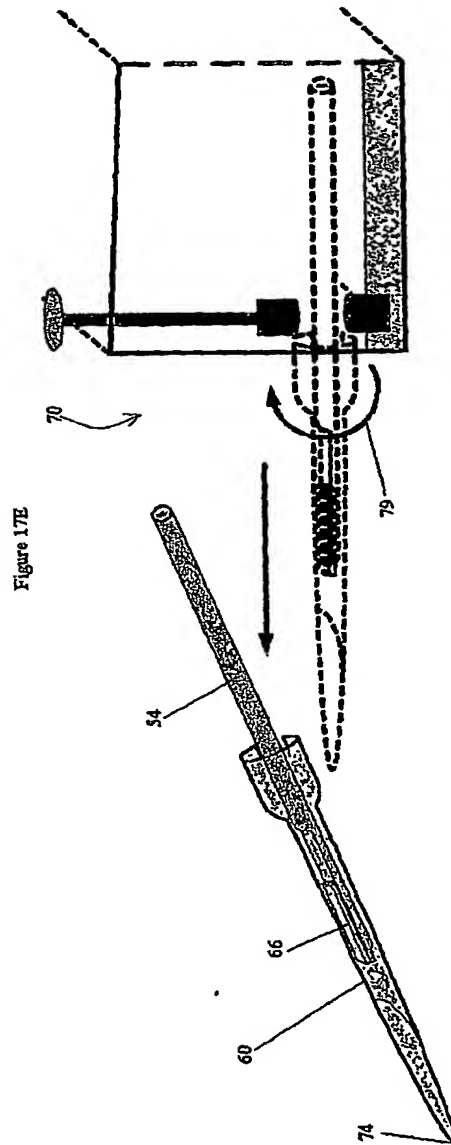






**Figure 17C**

Figure 17D  
58





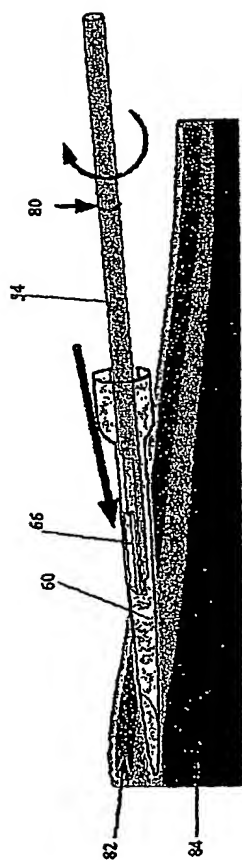


Figure 18A

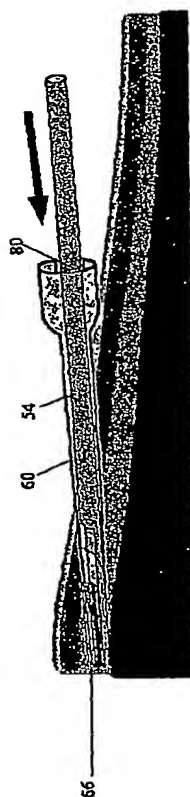


Figure 18B

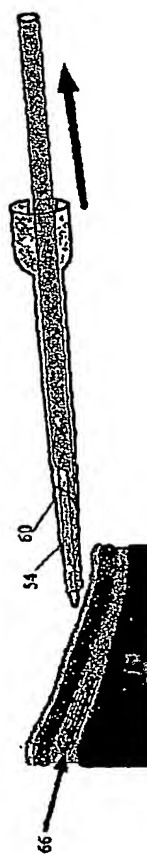


Figure 18C

Figure 19A



Figure 19B

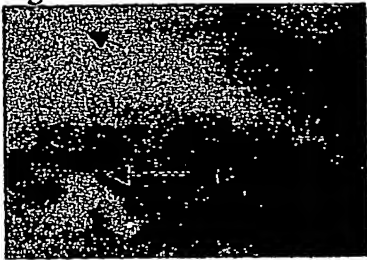


Figure 19C

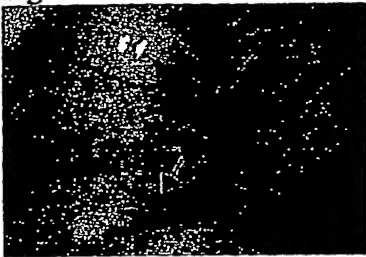


Figure 20a



Figure 20b

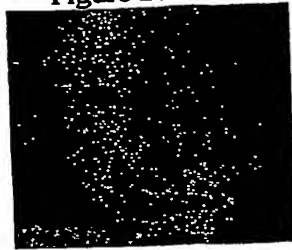


Figure 21A

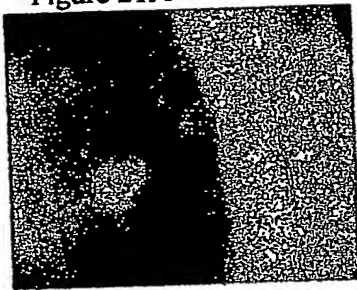


Figure 21B



Figure 22A

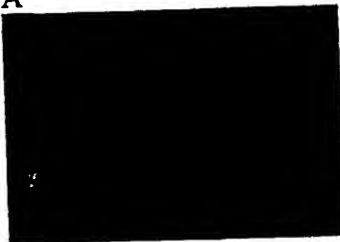


Figure 22B



Figures 23A-C

A



B



C



FIGS 23 D→G

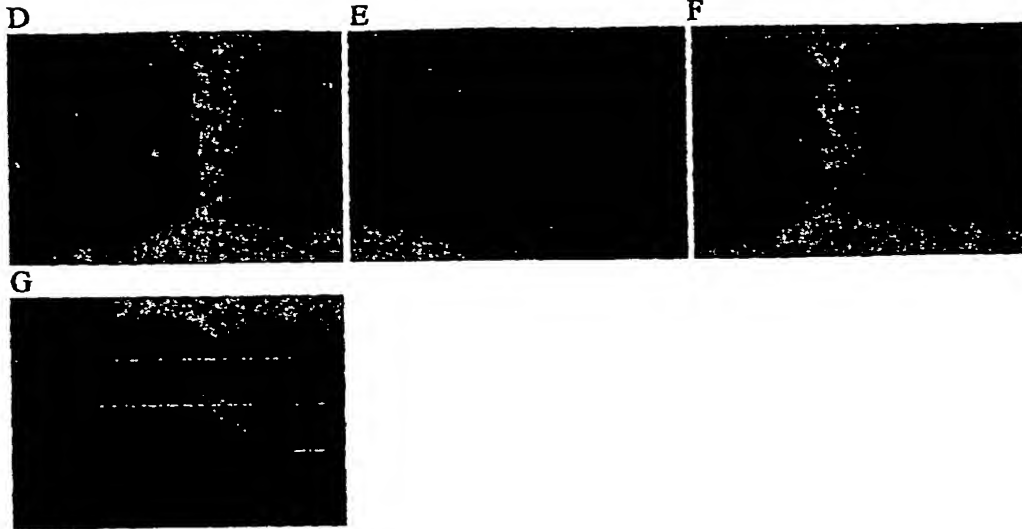
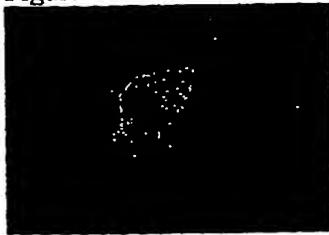


Figure 24



## SEQUENCE LISTING

<110> Eduardo N. Mitrani

<120> METHOD AND DEVICE FOR INDUCING BIOLOGICAL PROCESSES BY MICRO-ORGANS

<130> 23451

<160> 8

<170> PatentIn Ver. 3.1

<210> 1

<211> 21

<212> DNA

<213> Artificial sequence

<220>

<223> Description of Artificial sequence: PCR primer

<400> 1

CGTGGGTGGA GGAGGGTGA C 21

<210> 2

<211> 21

<212> DNA

<213> Artificial sequence

<220>

<223> Description of Artificial sequence: PCR primer

<400> 2

TGCGTCAAAC CACCAGCCTC C 21

<210> 3

<211> 20

<212> DNA

<213> Artificial sequence

<220>

<223> Description of Artificial sequence: PCR primer

<400> 3

TACCACAGGC ATTGTGATGG 20

<210> 4

<211> 18

<212> DNA

<213> Artificial sequence

<220>

<223> Description of Artificial sequence: PCR primer

<400> 4

AATAGTGATGA CCTGGCCGT 18

<210> 5

<211> 20

<212> DNA

<213> Artificial sequence

<220>

<223> Description of Artificial sequence: PCR primer

<400> 5

GGTCACACAG GGACAGCAGG 20

<210> 6

<211> 21

<212> DNA

<213> Artificial sequence

<220>

<223> Description of Artificial sequence: PCR primer

<400> 6

CCAAGGGCCG GATCAGCATG G 21

<210> 7

<211> 19

<212> DNA

<213> Artificial sequence

<220>

<223> Description of Artificial sequence: PCR primer

<400> 7

ACTTTCTGCT CTCTTGGGT 19

<210> 8

<211> 18

<212> DNA

<213> Artificial sequence

<220>

<223> Description of Artificial sequence: PCR primer

<400> 8

CCGCCCTGGC TTGTCACA 18